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# THE STRUCTURAL BASIS OF T CELL ACTIVATION BY SUPERANTIGENS

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#### **ABSTRACT**

Superantigens (SAGs) are a class of immunostimulatory and disease-causing proteins of bacterial or viral origin with the ability to activate large fractions (5-20%) of the T cell population. Activation requires simultaneous interaction of the SAG with the VA domain of the T cell receptor (TCR) and with major histocompatibility complex (MHC) class II molecules on the surface of an antigenpresenting cell. Recent advances in knowledge of the three-dimensional structure of bacterial SAGs, and of their complexes with MHC class II molecules and the TCR & chain, provide a framework for understanding the molecular basis of T cell activation by these potent mitogens. These structures along with those of TCRpeptide/MHC complexes reveal how SAGs oircumvent the normal mechanism for T cell activation by peptide/MIIC and how they stimulate T cells expressing TCR B chains from a number of different families, resulting in polyclonal T cell activation. The crystal structures also provide insights into the basis for the specificity of different SAGs for particular TCR & chains, and for the observed influence of the TCR of chain on SAG reactivity. These studies open the way to the design of SAG variants with altered binding properties for TCR and MHC for use as tools in dissecting structure-activity relationships in this system.

#### INTRODUCTION

Tlymphocytes recognize a wide variety of antigens through highly diverse cell-surface glycoproteins known as T cell receptors (TCRs). These disulfide-linked heterodimers are composed of  $\alpha$  and  $\beta$ , or  $\gamma$  and  $\delta$ , chains that have variable (V) and constant (C) regions that are structurally homologous to those of antibodies (1, 2). Unlike antibodies, however, which recognize antigen alone,  $\alpha\beta$  TCRs recognize antigen only in the form of peptides bound to major histocompatibility complex (MHC) molecules. In addition, TCRs interact with a class of viral or bacterial proteins known as superantigens (SAGs), which stimulate T cells bearing particular  $V\beta$  elements, resulting in the massive release of T cell-derived cytokines such as interleukin (IL)-2 and tumor necrosis factor (TNF)  $\beta$ , generally followed by the eventual disappearance or inactivation of responding T cells (3-4). Activation of the T cell requires simultaneous interaction of the SAG with the TCR and with MHC class II molecules on an antigen-presenting cell (APC).

The best-characterized group of SAGs belongs to the so-called pyrogenic toxin SAG family, which includes the staphylococcal enterotoxins (SE)A through I (except F), staphylococcal toxic shock syndrome toxin-1 (TSST-1), streptococcal superantigen (SSA), and streptococcal pyrogenic exotoxins (SPE)A-C and -F (6-8). These bacterial SAGs have in common the following characteristics: (a) they are among the most potent pyrogens known, (b) they are all capable of inducing a highly lethal toxic shock syndrome, and (c) they share a typical three-dimensional structure consisting of two domains, termed large and small. The small domain is a fi-barrel made up of two fi-sheets, whereas the large domain contains a \$-grasp motif, an \alpha-belix packed against a mixed B-sheet that connects the peripheral strands (9-16). Nevertheless, each of these molecules has unique biological properties and stimulates the proliferation of T cells with different VB regions. Among their biological effects, the staphylococcal enterotoxins are characterized by their ability to induce emesis and diarrhea, whereas TSST-1 lacks emetic activity. The streptococcal toxins do not cause enteric problems but they are associated with cardiotoxicity. The level of sequence homology between the pyrogenic toxins varies widely, and they can be divided into groups based on sequence similarities. The highest degree of homology is achieved by SEA, SED, and SEE (between 53-81%), followed by the group of SEB, the SECs, SPBA, and SSA, with 50-66% of sequence homology. All the rest, including SPEB, SPEC, SPEF, and TSST-1, have poor or no homology to any other toxin, or to each other.

Bacterial SAGs that do not belong to the pyrogenic toxin family include the staphylococcal exfoliative toxins (ET) A and B (17, 18), Mycoplasma arthritidis mitogen (MAM) (19), and Versinia pseudotuberculosis mitogen (20, 21).

Among superantigenic proteins of viral origin, only mouse mammary tumor virus (MMTV)-encoded SAGs have been defined in detail (22). It has been demonstrated that mouse Mis endogenous SAGs are encoded by MMTV proviral DNA that has been integrated into the germline, demonstrating a link between endogenous SAGs and infectious agents. Other reports have shown superantigenic activity by the rabies virus nucleocapsid protein (23) and by two human tumor viruses, cytomegalovirus (24) and Epstein-Barr virus (25). Recently, the envelope gene of an endogenous human retrovirus isolated from panereatic islets was shown to encode an MHC class II—dependent SAG specific for VB7 (26).

The past four years have witnessed a remarkable series of advances in knowledge of the three-dimensional structure of TCRs (27-32) and of their complexes with peptide/MHC (33-35) and SAGs (36, 37). In this review, we focus on TCR-SAG interactions and describe current understanding of the structural basis of T cell activation by SAGs. After giving an overview of the biological effects of bacterial and viral SAGs, we discuss the affinity and kinetics of TCR and MHC binding to these molecules. We then describe the three-dimensional structures of MHC-SAG and TCR  $\beta$  chain-SAG complexes. These structures, along with those of TCR-peptide/MHC complexes, reveal how SAGs circumvent the normal mechanism for T cell activation by peptide/MHC and how they stimulate T cells expressing TCR  $\beta$  chains from a number of different families, resulting in polyclonal T cell activation. Finally, we discuss the structural basis for the specificity of different SAGs for particular TCR  $\beta$  chains and for the observed influence of the TCR  $\alpha$  chain on SAG reactivity.

#### BIOLOGICAL EFFECTS OF SUPERANTIGENS

### T Cell Anergy and Deletion

The specificity of interaction of SAGs with the  $V\beta$  domain of the TCR has provided a unique opportunity to examine the fate of reactive T cells in vivo independently of functional assays. Such studies have revealed that responding T cells can proliferate, become nonresponsive (energy), or even die (deletion) (38, 39). In the in vivo recognition of endogenous SAGs, intrathymic deletion in  $V\beta$ -specific subsets occurs at the double-positive (CD4\*, CD8\*) stage of development, and deletion is correspondingly apparent in both the mature CD4\* and CD8\* subsets (40-42). In the case of exogenous SAGs, an early report showed that mice injected from birth with SEB virtually lack  $V\beta3^+$  and  $V\beta8^+$  mature thymocytes, giving the first formal demonstration that clonal deletion can accompany induced tolerance to a foreign antigen (43). Subsequent studies confirmed this report and showed that in adult mice, SEB-specific mature T cells can, after an initial expansion, be rendered anergic in both in vivo and in vitro

models (44, 45). Moreover, SEB-induced death of  $V\beta8^+$  cells is independent of an intertripymus, because it also occurs in adult thymeetomized animals (46).

The presence of a SAG in the MMTV genome can guarantee the existence of actively dividing populations of T and B cells through the ability of the SAG to stimulate T cells, and, thus, let the virus complete its replication cycle (22). This was confirmed using mice transgenic for the MMTV (C3H) sag gene (47), The SAG-mediated deletion of V\$14+ T cells during early life conferred resistance to infection to these mice. Similar results were observed for a different exogenous MMTV, called SW, where the corresponding SAG stimulates V\$6+ cells (48). Because MMTV infection occurs during the shaping of the immune repertoire in neonatal life and because infection is persistent, there is a tradual deletion of SAO-reactive T cells (49). Finally, there is no evidence for a SAG-independent pathway of MMTV transmission, and only MMTV with functional sag genes can be transmitted through milk (50, 51). Once the SAG is stably integrated into the mouse genome, it can be inherited by successive generations. When expressed endogenously, it causes deletion of cognate T cells and prevents a reinfection with the same strain of virus that produces the SAG (47, 48).

### Toxic Shock Syndrome and Food Poisoning

Toxic shock syndrome (TSS) is an acute, life-threatening intoxication characterized by high fever, hypotension, rash, multiorgan dysfunction, and cutaneous desquamation that is caused by staphylococcal or streptococcal pyrogenic toxins (6, 52-54). The interaction of the pyrogenic toxins with TCR and MHC activates both the T cell, for secretion of TNF\$\beta\$, IL-2, and y interferon, and the APC, for secretion of TNF\$\alpha\$ and IL-1. The resulting massive cytokine release is believed to be responsible for capillary leak and hypotension, and it is also likely to cause the crythematous rash in TSS patients (52, 54).

Staphylococcal enterotoxins are among the most common causes of food poisoning in humans. It has been suggested that the enterotoxic effects are directly related to their superantigenic activity, i.e. dependent on T cell stiraulation and probably caused by massive cytokine release (3, 6). However, some evidence suggests that the emetic and T cell proliferative activities of the toxins may be distinct (3, 55-57). In fact, the induction of emesis has been attributed to leukotriene or histamine release (58). It has been shown that SEB and SEA can rapidly cross an epithelial membrane in intact, fully functional form, thus gaining access to T cells. On the contrary, TSST-1, which lacks emetic activity, although able to transcytose epithelium, may be more easily destroyed by digestive enzymes in the stomach and intestine (59). Thus, the ability to cause enterotoxicity may be related to the resistance to digestion of the enterotoxins.

#### Autoimmune Diseases

In eurojmmune diseases, a breakdown of self-tolerance leads to the generation of an immune response against a specific target antigen or antigens. A large body of clinical and epidemiological evidence indicates that infections are important in the induction of autoimmune disorders such as autoimmune myocarditis (60) and insulin-dependent diabetes mellitus (61). One mechanism by which this has long been thought to occur is through the activation of autoreactive T cells by epitones on microbial antigens that are cross-reactive with antigens on target organs (62). For example, a number of viral and bacterial poptides have been identified that efficiently activate myelin basic protein (MBP)-specific T cell clones from multiple sclerosls (MS) patients (63, 64). More recently, it has been proposed that SAGs derived from bacteria, mycoplasma, or viruses may initiate autoinnume disease by activating specific auti-self T cell clones (3, 65). Indeed, microbial SAGs have been shown to trigger clinical relapses of autoimmune disease in several animal models, as discussed below. The expansion of selected VB families in the affected organs or peripheral blood of certain individuals with autoimmune disease has also been documented (3,65). However, it is a common observation that different TCR repertoire studies of the same disease can provide different results (66). Even when it is unlikely that SAGs by themselves initiate an autoimmume disease (65), they may modulate disease pathogenesis. In susceptible individuals, the activation of autoreactive T cells is a necessary, but not sufficient, condition for the development of an autoimmune disease. A sufficient degree of clonal expansion of autoreactive T cells may be a major limiting factor, and SAGs may induce such an expansion. Alternatively, the activation of B lymphocytes and other APC through the SAG bridge may lead to the secretion of autoantibodies and interleukins that contribute to inflammation.

Evidence for an autoimmune origin of MS comes from (a) the presence of CD4+T cells and cells expressing MHC class II molecules in inflamed tissues (67), (b) the finding that MS is associated with certain MHC class II alleles (68), and (c) the demonstration that MBP-specific T cells are clonally expanded in MS patients (69-72). In experimental autoimmune encephalomyelitis (EAE), a model for MS, administration of SBB to PL/I mice following immunization with a peptide derived from MBP (Ac1-11) was found to induce paralysis in mice with subclinical disease and to trigger relapses in mice that are in remission following an initial episods of paralysis (73-75). It was shown that these effects are the direct result of stimulation by SEB of V\$8-expressing encephalitogenic T cells specific for MBP Ac1-11.

An analysis of the TCR  $\beta$  chain repertoire of synovial T cells from rheumatoid arthritis (RA) patients revealed a selective expansion of  $V\beta 14$ -bearing T cells compared with the levels in the peripheral blood of the same

individuals (76). A mechanism for the pathogenesis of RA was proposed in which a microbial SAG activates disease-mediating VB14+ T cells and allows these activated cells to enter the synovial tissue, where they persist because of reactivation by autoantigens. In collagen II-induced arthritis (CIA), a model for RA, mice are immunized with native porcine type II collagen and develop joint swelling. It has been demonstrated that T cells expressing VB8 are important in the development of CIA (77-80) and that administration of SEB 10 days prior to a collagen II challenge protects mice from CIA (81). In both EAE and CIA, the response to self-antigens is controlled by a potent regulatory T cell circuitry based on recognition of different determinants derived from the TCR V88.2 chain (81-83). The SAG MAM, which derives from a naturally occurring mouse arthritogenic mycoplasma, activates V 884 T cells. Administration of MAM has been shown to markedly exacerbate arthritis in mice that were convalescent from CIA, or to trigger arthritis in animals previously immunized with collagen II but that had failed to develop clinical disease (19).

Insulin-dependent diabetes mellinis (IDDM) is an autoimmune disease affeeting pancreatic & cells that scorete insulin. A relationship between viral infections and the development of IDDM has been long suspected (84). An analysis of pancreatic islet-infiltrating T cells from patients with IDDM revealed preferential expression of the V\$7 gene segment, but no selection for particular Va segments or VB-D-JB junctional sequences (85, 86). This led to the proposal that a SAG associated with pancreatic isless may be involved in the pathogenesis of IDDM. This putative SAG was recently identified by Conrad et al (26), who isolated a novel human endogenous retrovirus from supernatants of IDDM islets and showed that the envelope gene encodes an MHC class II-dependent SAG specific for VB7.

Kawasaki disease (KD) is an acute febrile illness with symptoms similar to toxic shock syndrome. Several studies have revealed a significantly elevated level of circulating V\$2+ and, to a lesser extent, V\$8.1+ T cells in patients with acute KD, compared with control populations (87-90). Sequencing of these & chains revealed extensive junctional region diversity, which suggests activation by SAG and not a specific disease-associated antigen. Bacteriaproducing toxins that activate VB2+ T cells (TSST-1 and SPEB/SPEC) were isolated from 13 out of 16 KD patients but only 1 out of 15 in the control group (91). Nevertheless, other groups were not able to document the expansion of any Vß family during the scute phase of KD (92,93). Differences in population studied, method and time of collecting samples, and techniques used could potentially explain the differences in results. Alternatively, the expansion of selected VA families may not be related to the pathogenesis of the disease (94).

#### Skin Diseases

The staphylococcal toxins ETA and ETB induce the symptoms associated with staphylococcal scalded skin syndrome, characterized by a specific intraepidermal separation of layers of the skin (95). X-ray crystallographic studies of ETA have shown that its overall structure is similar to that of the chymotrypsin-like serine protease family of enzymes (17.18). The catalytic triad includes the residue Ser195, which when mutated to cysteine abolishes the characteristic separation of epidermal layers, although the ability to induce T cell proliferation is not altered (17.96). These findings suggest that skin separation is the result of a specific proteolysis by ETA, and not related to its superantigenic activity; the latter is probably involved in the edema or redness associated with scalded skin syndrome.

Psoriasis is a disease characterized by increased proliferation of epidermal cells associated with an inflammatory component. Patients with acute gutate psoriasis often have flares of psoriasis following streptococcal infections (97). Histological examination of early skin lesions shows that infiltration of lymphocytes and macrophages into the skin precedes the characteristic epidermal proliferation of psoriasis. The predominant distribution of  $V\beta 2$ -,  $V\beta 3$ -, and  $V\beta 5$ -bearing T cells in lesional skin of acute gutate psoriasis has been described (98-100). However, there are conflicting reports about the restricted T cell receptor repentoire in chronic psoriasis (98, 101), and no increase in SAG-producing Staphylococcus ameus has been seen in chronic psoriatic patients (102). It is unlikely, then, that SAGs are essential to the continuance of psoriasis, although they may be exacerbating factors or triggers for the disease.

Atopic dermatitis is a chronic pruride inflammation of the skin characterized by local infiltration of monocytes and lymphocytes, mast cell degranulation, and immediate and delayed hypersensitivity (103). There are numerous reports that S. aureus can exacerbate this disease, and S. aureus was isolated from the affected skin of more than 90% of patients (104). More than half of the patients had S. aureus that secreted SEA, SEB, and TSST-1 (105). Sera from 57% of atopic dermatitis patients contained immunoglobulin E specific for one or more of these SAGs. Thus, epicutaneous superantigenic toxins might induce specific immunoglobulin E in atopic dermatitis patients, as well as mast cell degranulation.

## AFFINITY AND KINETICS OF TCR BINDING TO SUPERANTIGENS

Gascoigne & Ames first demonstrated direct binding of a soluble TCR  $\beta$  chain (mouse  $V\beta$ 3) to SEA presented by MHC class II molecules on cells (106).

However, the affinity was too low to be measured in their cell-binding assay. More recently, the development of surface plasmon resonance techniques for detecting macromolecular interactions (107) has permitted the precise measurement of kinetic and affinity constants for TCR binding to SAGs, as well as to peptide/MHC complexes (108). By this method, a soluble human TCR (V/3.1) was found to bind immobilized SEB in the absence of MHC class II molecules with a dissociation constant  $(X_D)$  of 0.8  $\mu$ M; the on-rate  $(k_{on})$  of the interaction was  $1.3 \times 10^6 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$  and the off-rate  $(k_{\rm eff}) \,1.1 \times 10^{-2} \,\mathrm{s}^{-1}$  (109). Specific binding of soluble 14.3.d TCR \$\beta\$ chain (mouse V\$8.2) was demonstrated to SEB, SEC1, SEC2, SEC3, and SPEA, consistent with the known proliferative effects of these SAGs on T cells expressing V\$8.2 (110). In contrast, SBA, which does not stimulate V\$8.2-bearing cells, did not bind the recombinant B chain. Affinities ranged from 3 µM for SEC3 to 140 µM for SEB; kan and  $k_{\rm eff}$  were too fast to be accurately measured, but were estimated at > 105 M<sup>-1</sup> s<sup>-1</sup> and >0.1 s<sup>-1</sup>, respectively (110, 111). The unpaired  $\beta$  chain was shown to fully retain the SAG-binding activity of the assembled 14.3.d af TCR heterodimer (110). A KD of 1.1 µM was measured for the binding of SEC2 to the mouse D10 TCR (V#8.2), with a  $k_{on}$  of 1.7 × 10° M<sup>-1</sup> s<sup>-1</sup> and a  $k_{on}$  of 1.9 × 10<sup>-2</sup> s<sup>-1</sup> (112). These values closely resemble those for the interaction of D10 TCR with its cognate popular/MHC class II ligand, which has a  $K_D$  of 2.1  $\mu M$  and a  $k_{on}$ and  $k_{\text{off}}$  of 1.0  $\times$  10<sup>4</sup> M<sup>-1</sup> s<sup>-1</sup> and 2.2  $\times$  10<sup>-2</sup> s<sup>-1</sup>, respectively (112).

In each of the above examples, TCR-SAG binding is characterized by low affinities (>10<sup>-6</sup> M) and very fast  $k_{on}$  and  $k_{off}$  (>10<sup>4</sup> M<sup>-1</sup> s<sup>-1</sup> and >10<sup>-2</sup> s<sup>-1</sup>, respectively). It is noteworthy that low affinities and rapid dissociation kinetics have also been reported for the interaction of other T cell surface glycoproteins with their ligands, such as the adhesion molecule CD2 with CD48 (113, 114). In particular, the affinities of TCR-SAG interactions (10-4-10-6 M) are remarkably similar to those reported for the binding of TCRs to specific peptide/MHC complexes (10<sup>-4</sup>-10<sup>-7</sup> M) (115-117) and are much weaker than those of antigen-antibody reactions (typically 10-8-10-11 M). In the case of adhesion molecules, fast dissociation rates may facilitate deadhesion, a requirement for cell motility (113). In the case of TCRs, rapid off-rates may permit a single peptide/MHC complex to sequentially bind and trigger a large number of TCRs (up to 200), as proposed in the social triggering (118, 119) and kinetic prooficading (120) models of T cell activation, until a certain activation threshold is reached. The finding that the binding of bacterial SAGs to the TCR is characterized by low affinities and fast dissociation kinetics suggests that SAGs mimic the interaction of peptide/MHC complexes with the TCR in terms of affinity and kinetics and that some form of serial engagement may also operate in T cell activation by SAGs (111, 119, 121).

The relationship between the affinity of SAGs for TCR and MHC and their ability to activate T cells has been investigated using site-directed mutants of SEC3 and SEB (111). In order to mimic normal physiological conditions as closely as possible, resting lymph node T cells bearing the 14.3.d TCR from RAG-2"- TCR transgenic mice were used to measure the stimulatory effects of mutant SAGs on BALB/c spleen cells expressing I-Ed, or on MHC class II-negative mouse fibroblasts transfected with a gene encoding human leukocyte antigen (HLA)-DR1. A clear and simple relationship was observed between the affinity of SAGs for the TCR and their mitogenic potency: the tighter the binding of a particular mutant of SEC3 or SEB to the TCR ff chain, the greater its ability to stimulate T colls. The affinities of the SAGs tested ranged from 3.5 µM to >250 µM. However, an apparent exception to this simple affinity-activity rule was the finding that SEB stimulated transgenic T cells about 10-fold botter than did SEC3, even though the affinity of SEB for the TCR & chain (140  $\mu$ M) is much lower than that of SEC3 (111). To determine whether the surprisingly strong mitogenic potency of SEB relative to SEC3 could be attributed to tighter binding to MHC class II on AFC, the binding of SEB and SEC3 to soluble recombinant HLA-DR1 was measured by sedimentation equilibrium: Whereas SBB bound to DRI with a KD of 14 µM, the corresponding value for SEC3 was 48 µM. Therefore, the unexpectedly high mitogenic potency of SEB relative to SEC3 can be explained by the stronger binding of SEB to MHC class II. This indicates that mitogenic potency is the result of an interplay between TCR-SAG and SAG-MHC interactions, such that a relatively small (threefold) increase in the affinity of a SAG for MHC can overcome a large (35-fold) decrease in the affinity of a SAG for the TCR.

With the apparent affinities of SAGs for both TCR and MHC class II molecules in the micromolar range, nearly all SAG molecules will be unbound at physiological SAG concentrations (10-12-10-15 M) (121). Under these conditions, it is difficult to understand how a SAG can effectively cross-link the T cell and APC. The problem is seemingly less severe for peptide/MHC because the pentide is, in effect, irreversibly bound to MHC. One possible explanation for the ability of SAGs to trigger T cells at concentrations orders of magnitude less than their Kos is that accessory molecules such as CD4 help stabilize the TCR-SAG-MHC complex sufficiently for activation to occur. Another is that the overall stability of the TCR-SAG-MHC complex is considerably greater than would be expected from considering the TCR-SAG and SAG-MHC interactions independently. That is, the binding of SAGs may be a cooperative process in which the SAG-MHC complex binds the TCR with greater affinity than does the SAG alone. This hypothesis is supported by the finding that the affinity of SEB for a soluble human TCR was significantly enhanced by the addition of soluble HLA-DR1 (109). The potential role of the TCR a chain in stabilizing the TCR-SAG-MHC complex is discussed in a later section.

## STRUCTURE OF SUPERANTIGEN-MHC CLASS II COMPLEXES

The three-dimensional structures of three SAG-MHC class II complexes have heen determined by Wiley and colleagues: (o) the complex between SEB and HLA-DRI to 2.7-A resolution (122), (b) the complex between SEB and HLA-DR4 to 2.5-Å resolution (123), and (c) the complex between TSST-1 and HLA-DR1 to 3.5-Å resolution (124). In the SEB-HLA-DR1 complex (Figure 1.4) (see color plates)], SEB binds to the all domain of DR1, contacting residues from the first and third turns of the \$-sheet and from the N-terminal portion of the  $\alpha$ -helix (122). The binding of SEB to DR4 is similar (123). The ability of SEB to bind many different DR alleles can therefore be explained by its exclusive interaction with the DRI & chain, which is conserved in all DR molecules. Residues of SEB in contact with DR1 derive mainly from the small domain of the SAG, although several residues from the large domain also contact the DR & chain. SEB binds away from the peptide-binding groove of DRI and does not contact the bound peptide. The affinity of SEB for DRI was reported as approximately 0.5 12M in a cell-binding assay (125) and as 14 µM using soluble DR1 (111).

Although the TSST-1 binding site on HLA-DR1 overlaps that of SBB, the two SAGs bind differently (124). Whereas SBB binds primarily off one edge of the peptide-binding groove (Figure 1A), TSST-1 extends over nearly half the binding groove and contacts the  $\alpha$ -helix of the  $\alpha$ 1 domain of DR1, the bound peptide, and part of the  $\alpha$ -helix of the  $\beta$ 1 domain of DR1 (Figure 1B). This binding mode suggests that the interaction of TSST-1 with MHC class II molecules may be partially peptide dependent. In agreement with the crystal structure, certain peptides were found to promote the presentation of TSST-1 by I-Ab up to 5000-fold (126). In contast, the binding of SBB to I-Ab and I-Edc is peptide independent (126, 127).

Although no crystal structures have been reported for SEA complexed with MHC class II molecules, mutagenesis and binding studies have demonstrated that SEA possesses two distinct, yet cooperative, binding sites for class II molecules: (a) a low-affinity site ( $K_D = 10^{-5}$  M) to the DR1  $\alpha$  chain analogous to the DR1-binding site of SEB, and (b) a  $Z_0^{2d}$ -dependent, high-affinity site ( $K_D = 10^{-7}$  M) to the polymorphic DR1  $\beta$  chain (127–129). Binding of one SEA molecule to the DR1  $\beta$  chain enhances the binding of a second SEA molecule to the DR1  $\alpha$  chain (128, 129). Supprisingly, mutations in the  $Z_0^{2d}$ -dependent site completely abolish SEA activity, even though it can still bind the DR1  $\alpha$  chain through its low-affinity SEB-like site. This suggests that MHC cross-linking on the surface of APC may be an essential feature of SEA function. This conclusion is supported by the demonstration that SEA<sub>2</sub>-DR1

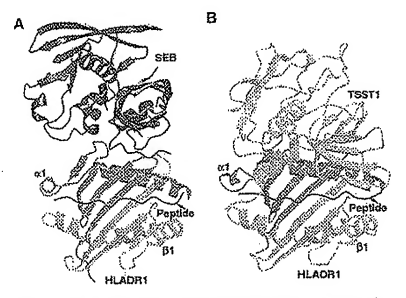


Figure / Timeo-dimensional structures of the complexes between HLA-DR1 and SEB and between HLA-DR1 and TSST-1. (4) Ribbons diagram of the SEB-HLA-Dr1 complex (122). (8) Ribbons diagram of the TSST-HLA-DR-1 complex (124). Colors are as follows: SEB (blue), TSST-1 (pink), DR1 of domain (green), DR1 \$ domain (yellow), and peptide (red).

trimers exist in solution (130), as well as by the finding that SEA mediates signaling through the APC by direct cross-linking of DR1 molecules (131, 132). The high sequence identity between SEE and SEA (see Figure 3A) suggests that they may bind class II molecules similarly. Two additional SAGs have been described that cross-link class II molecules, but by different mechanisms than SEA. The crystal structure of SED shows that this SAG forms dimers in the presence of Zn2+ by coordinating two Zn2+ ions between the large domains of two SED molecules (15). Binding to MHC class II molecules is believed to occur through the small domain to the class II a chain in a manner similar to SEB, resulting in a tetrameric class II α-SED-SED-class II α complex on the APC. The three-dimensional structure of SPEC reveals that the class II a chainbinding site on the small domain has been replaced by SPEC dimer interface (16). Instead, SPEC binds only to the class II & chain. This could potentially lead to the formation of class II B-SPEC-SPEC-class II B tetramers. Dimeric SAGE like SED and SPEC may facilitate TCR dimerization and subsequent T cell triggering.

Endogenous SAGs encoded by MMTV can be efficiently presented to T cells only by B cells, through interaction with MHC class II molecules (22). Although direct binding has been demonstrated between recombinant forms of MMTV SAGs and MHC class II molecules (133, 134), the interaction remains poorly understood. I-E molecules are the best presenters for all the described MMTV SAGs (135). In addition, C57BL mice that lack I-E molecule, and thus are not able to present SAG to T cells, are resistant to milk-borne MMTV (C3H) (136). Analysis of class II mutants that lost the ability to present bacterial SAGs revealed that bacterial SAGs have different binding requirements than do MMTV SAGs (137). Another study showed, however, an overlap in at least one binding site for MMTV and SEA on the MHC molecule (138). Recently it was shown that N-linked glycosylation is required for effective B cell presentation of MMTV SAGs to T cells (139).

# STRUCTURE OF TCR $\beta$ CHAIN-SUPERANTIGEN COMPLEXES

The three-dimensional structures of several TCR  $\beta$  chain-SAG complexes have been determined to date, each involving the  $V\beta C\beta$  chain of the mouse 14.3.d TCR specific for a hemagglutinin peptide of influenza virus bound by the I-E<sup>d</sup> class II molecule: (a) the complex between the  $\beta$  chain and SEC2 to 3.5-Å resolution (36), (b) the complex with SEC3 to 3.5-Å resolution (36), (c) the complex with SEB to 2.4-Å resolution (37), and (d) the complex with a mutant of SBB in which value at position 26 is replaced by tyrosine (SEB V26Y) to 2.6-Å resolution (37). The SEB V26Y mutant was designed on the

basis of the structure of the TCR  $\beta$ -SEC3 complex to bind the  $\beta$  chain more tightly than wild-type SEB: Its  $K_D$  is 12  $\mu$ M, approximately 12 times lower than that of SEB ( $K_D=140~\mu$ M), but still four times higher than that of SEC3 ( $K_D=3~\mu$ M) (111). The mutant is fourfold more active in T cell proliferation assays than is wild-type SEB, consistent with its enhanced affinity.

The crystal structures of the TCR  $\beta$ -SEC2/3 complexes identified the regions of the  $\beta$  chain recognized by SEC and showed how SAGs circumvent the normal mechanism for T cell activation by specific peptide/MHC complexes. However, the moderate resolution of these structures (3.5 Å) precluded a detailed analysis of the interface between the two proteins in terms of hydrogen bonds, van der Waals interactions, and solvent structure. To achieve a more complete description of a  $\beta$ -SAG interface, as well as to assess whether conformational changes occur in either or both proteins upon complex formation, the structures of the complexes between the 14.3.d  $\beta$  chain and SEB and SEB V26Y were determined to high resolution (37). These structures, along with those of  $\alpha\beta$  TCR heterodimers (29-31) and TCR-peptide/MHC complexes (33-35), can account for the specificity of different SAGs for particular  $\beta$  chains and for the influence of the TCR  $\beta$  chain on SAG reactivity (5, 140-142).

## Overall Structure of the TCR $\beta$ -SEB and TCR $\beta$ -SEC Complexes

The overall structure of the  $\beta$ -SEB complex is shown in Figure 2A (see color plates). The complex is formed through contacts between the V $\beta$  domain and the small and large domains of SEB. The complementarity-determining region (CDR)2 of the TCR  $\beta$  chain and, to lesser extents, hypervariable region (HV)4 and framework regions (FR)2 and -3 bind in the cleft between the two domains of the SAG (37). This binding mode is similar to that observed in the  $\beta$ -SEC2 and  $\beta$ -SEC3 complexes (36), but with several differences, as discussed below. SEC2 and SEC3 bind identically to the 14.3 d  $\beta$  chain, and none of the four amino acid differences between SEC2 and SEC3 is located in the complex interface. This is reflected in the  $K_D$ s of the two SAGs, which are both approximately 3  $\mu$ M. The TCR-binding sites of SEC3 and SEB, in contrast, differ at positions 20, 26, and 91; these differences presumably account for the 45-fold weaker affinity of SEB for the 14.3 d  $\beta$  chain (111).

The solvent-excluded surface area for the  $\beta$ -SEB complex is 1343 Å<sup>2</sup> (685 Å<sup>2</sup> from V $\beta$  and 658 Å<sup>2</sup> from SEB); the buried surface area for the  $\beta$ -SEC3 complex is similar (1300 Å<sup>2</sup>). These values are within the range observed for antigen-antibody complexes (143) but somewhat smaller than the approximately 1800 Å<sup>2</sup> of buried surface in TCR-peptide/MHC complexes (33–35). As shown in Figure 24, the TCR-binding site of SEB is adjacent to, but distinct from, the MHC-binding site of this SAG (122, 123). This spatial proximity

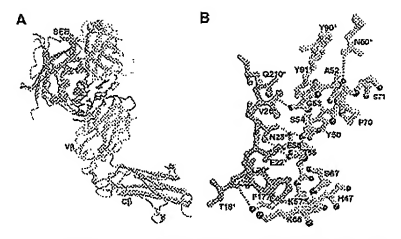


Figure 2 TCR \$\text{SEB camplex.} (A) Ribbeas diagram of the V\$C\$\theta\$-SEB camplex (37). Colors are as follows: V\$\theta\$ (yellow), CDR1 (pink), CDR2 (red), CDR3 (grap), HV4 (blue), C\$\theta\$ (brown), SEB large domain (green), and SEB small domain (blue). Residues of V\$\theta\$ and SEB involved in interactions in the TCR-SAG interface are red. Residues of SEB in contact with MHC in the attuature of the SEB-HLA-DR1 complex (122) are yellow. The SEB distulfide loop (light grap), which is not visible in the electron density map of the \$\theta\$-SEB complex, was modeled according to the uncomplexed SEB crystal structure (9). (B) Interactions in the \$\theta\$-SEB interface. View is the same as in ponel A. V\$\theta\$ atoms are colored accordingly to atom type; carbon, altrogen and exygen atoms are yellow blue, and rad, respectively. SEB clams are colored green (large domain) and blue (small domain). SEB residues are indicated with asterisks. Hydrogen bonds are dotted brown lines.

suggests that the two binding sites may not be completely independent; that is, the affinity of the TCR for SEB alone may be lower (or higher) than its affinity for SEB bound to MHC class II molecules.

### Structure of the B-SEB and B-SEC Interfaces

The V\$ residues in contact with SEB are as follows: His47 of FR2; Tyr50. Alasz, Glysz, Serze, and Three of CDR2; Gluse, Lyser, Tyres, Lysee, and Ala67 of FR3; and Pro70 and Scr71 of HV4 (Table 1). The FR2, CDR2, FR3, and HV4 regions contribute 7%, 50%, 34%, and 9%, respectively, of the total contacts to SEB. The crystal structure therefore readily accounts for mutational and genetic evidence implicating VB CDR2 and HV4 in SAG recognition (3). In the \$-SEC3 complex (36), the V\$ residues in contact with the SAG are as follows: Tyr50, Ala52, Gly53, Ser54, and Thr55 of CDR2; Glu56, Lys57, and Lys66 of FR3; and Pro70 and Ser71 of HV4 (Table 1). The CDR2, FR3, and HV4 regions contribute 63%, 32%, and 7%, respectively, of the total contacts to SEC3. Thus, although there are several differences in contacting residues in the two complexes (e.g. V# FR2 His47, which contacts SEB but not SEC3, and V/S CDR2 Gly51, which contacts SEC3 but not SEB), CDR2 and FR3 account for the majority of intersections with the SAG in both complexes, with HV4 playing only a secondary role. The binding sites on the TCR for SAG and peptide/MHC class I molecules only partially overlap. As shown in Table 1, only V\$ residues Tyr50, AlaS2, Thr55, and Glu56 contact both SEB and peptide/MHC in the 2C TCR-dEV8/H-2Kb complex (34).

The SAG residues in contact with  $V\beta$  are as follows: Asn60, Tyr90, and Tyr91 (Val91 in SEC3) of the small domain; and Tur18, Gly19, Lcu20 (Thr20 in SEC3), Glu22 (in  $\beta$ -SEB only), Asn23, Tyr26 (in  $\beta$ -SEC3 and  $\beta$ -SEB V26Y only), Phe177 (Phc176 in SEB), and Glu210 of the large domain (Table 1). Residues Asn23, Asn60, and Tyr90 are strictly conserved among bacterial SAGs reactive with mouse  $V\beta$ 8.2, including SEC1-3 and SPEA, and have been shown to constitute energetic hot spots for binding the 14.3.d  $\beta$  chain (111) (Figure 3A, see color).

The structures of the  $\beta$ -SEB and  $\beta$ -SEC complexes enable us to understand how SEB and SEC, which have nearly identical V $\beta$  specificities, can each stimulate T cells expressing V $\beta$  domains from a number of different families (3). As shown in Figure 2B, all the hydrogen bonds between SEB and V $\beta$  are formed between SEB side-chain atoms and V $\beta$  main-chain atoms, except for a hydrogen bond between the main-chain oxygen of SEB Thr18 and the side chain of V $\beta$  Lys57 (SEB Thr18 O-N $\zeta$  Lys57 V $\beta$ ). Four of the main-chain-side-chain hydrogen bonds in the  $\beta$ -SEB complex are also present in the  $\beta$ -SEC3 complex: V $\beta$  Gly53 O-N $\epsilon$ 2 Gln210 SEB, V $\beta$  Thr55 N-O $\epsilon$ 1 Asn23 SEB, V $\beta$  Thr55 O-N $\epsilon$ 2 Asn23 SEB, and V $\beta$  Pro70 O-N $\epsilon$ 2 Asn60 SEB (Table 1).

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eng 2	REGP	DP TP	DELN	KAFR	-PT C	LXB	HOCV L	מ-פצ	107 V	AYKV	X2AD	KTLK	HDLI	YHISDKK
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SPGR	-200	DP DP	SQLK	P.Z\$ -	LVK	mre	IXXI	YTE - C	wy T	MEIN	KZVD	QLL5	X.VdK	assvay
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Figure 3 Residues defining the interection of bacterial SAGs with TCR  $\beta$  chains. (4) Sequence nligament of bacterial SAGs (SEB, SEC1-3, SPEA, SEA, SED, SEE, SPEC, TSST-1) based on structural information. The  $\alpha$ -carbon skeletows were first optimally superposed. Sequences were then manually adjusted to minimize the number of gaps while respecting the structural skullenty. Residues 116-173 and 232-259 are not shown. SEC3 residues in contact with the TCR  $\beta$  chain are boxed in colors according to the loss of hinding free energy ( $\Delta$   $\Delta$ G) upon attaine substitution: (rad) > 2.5 kcallenol; (green) 0.5-1.5 kcallenol; (blue) < 0.5 kcallenol (111). The homologous residues in the other SAGs are only based in color if they are identical with those in SEC3. SEB residues contacting MHC in the crystal structure of the SEB-HLA-DR1 complex (121), and the corresponding residues of SEC3, are boxed in color if identical in SEB and SEC3 and in magento if different. (B) Sequence alignment of selected mouse (m) and human (h) TCR  $\beta$  chains reactive with SEB or SEC(3). Only SAG-containing residues no shown.

Table 1 Contacts between the 14.3.4 TCR & chain and stophylocoreal SAGe\*

		Hydroge	a bonds				
β		SE		2BC3p			
GS3	0	Q210	Ne2	Q2	0 Nr2		
7\$5°	N	N23	081	N2:			
	0	•	N82		78%		
	Ò			17.0	Oyl		
<b>£57</b>	NE	TIE	0				
P70	0	NGO	N82	1760	37,52		
***************************************	V	an des W	क्योर देशक	nck <sub>q</sub>			
******	<del>,</del>	No.	of		No. of		
ß	SEB	contacts		SECI	contact		
1147	L.7.0	~~~~~	1				
	F177		4				
YS0°	Y91	1	9	V93	1		
GSI				V91	4		
AS2º	Y90		4	Y90	5		
G\$3	3723		1	N23	1		
				Y26	7		
				Q210	4		
\$54	1552		2	N23	5		
				<b>Y91</b>	į		
1'55°	L20		4	1.50	2		
	N23		2	N23	4		
	E22		2				
	F177		1				
£56°	L.20		1	T20	1		
	N23		l				
K57	G19		3	GIY	4		
	L20		2	T20	4		
Y65	F177		1				
K66	· F177		6	F176	7		
A67	F177		4	F176	3		
P70	N60		2	1,58	1		
\$71	N60		2	<b>3</b> 460			

<sup>\*</sup>TCR, T cell inceptor: SAG, superunigen; SE, staphylococcal enterotoxib; MITC, melor litateoempetibility complex.

\*Data for the \$SEB and \$SEC3 complexes are from itelerances
37 and 36, respectively.

\*V\$ residues in contact with poption/AIHC in the 2C TCR-\$EV&II.
2K8 complex (34).

\*Van dez Waals contacts <4.0 Å.

complex formation, as also noted for the B-SEC3 complex (36). However, a number of adjustments in VB and SAG side-chain positions are evident when comparing the free and bound structures of the 14.3.d  $\beta$  chain and SEB. Certain of these changes are necessary to avoid storic clashes between the TCR and SAG, whereas others probably serve to maximize productive interactions between the two proteins. For example, the side chain of SEB residue Tyr91 undergoes a 120° rotation away from VB in order to avoid a collision with CDR2 (37). The structural rearrangements in the B-SEB interface are of similar magnitude to those observed in antigen-antibody complexes in which the antigen is a protein (146, 147, 152, 153). They imply a limited "induced fir" mechanism for TCR-SAG recognition analogous to that described for antigen-antibody interactions (152, 154). A further indication of flexibility in TCR-SAG association comes from the finding that the two  $\beta$ -SEB molecules in the asymmetric unit of the crystal, although similar, are not identical. A rotation of 6° is required to optimize the overlap between SEB molecules in the two complexes following superposition of their  $V\beta$  domains (37).

The changes in the conformation of interface residues in the 14.3.d B-SEB complex are not as large as those in the 2C TCR-dEV8/H-2Kb complex, in which the CDR1 and CDR3 loops of the Va domain are displaced 4-6 Å reiztive to their positions in the unliganded 2C TCR structure (34). This probably indicates that the SAG has evolved to optimize its fit to the TCR. Indeed, calculations of shape complementarity (155) reveal that the  $\beta$ -SEB interface is about as tightly packed as antigen-antibody interfaces but significantly more tightly packed than TCR-peptide/MHC class I interfaces (33, 34, 156). It is important to emphasize, however, that, for both 2C TCR-dEV8/H-2K6 and 14.3.d TCR  $\beta$ -SEB complexes, the observed conformational changes are localized to the interfaces between the proteins and are not transmitted to the constant regions of the TCR. Thus, the possibility that changes in TCR conformation upon ligand binding are responsible for initiating T cell signaling can probably be ruled out. Rather, mechanisms based on ligand-induced TCR oligomerization (157) are more likely to account for T cell activation by peptide/MIIC or SAGs.

## Structural Basis for the V\$-binding Specificity of SEB and SEC

The structure of the 14.3.d  $\beta$  chain-SEB complex explains why SEB recognizes certain V $\beta$  families but not others. As discussed above, all the hydrogen bonds between SEB and mouse V $\beta$ 8.2 are formed between SEB side chains and V $\beta$  main-chain atoms (Figure 2B, Table 1), such that the positions of these main-chain atoms should be similar in V $\beta$  domains reactive with SEB but significantly different in V $\beta$ 8 that do not bind this SAG. A comparison of V $\beta$ 8 domains

of known three-dimensional structure confirms this expectation. Thus, SEB activates T cells bearing mouse VB8 and human VB12, but not mouse VB2 or VBS (3). When human VB12.3 (33) is superposed onto mouse VB8.2 (Figure 4A, see color plates), the RMS difference in  $\alpha$ -carbon positions for 14 residues in the SEB-binding site is only 0.9 Å. However, when mouse V\$2.3 (30) is superposed onto mouse V\$8.2 (Figure 4B), the RMS difference is 3.0 Å. This difference is largely attributable to a strand switch in  $V\beta 2.3$  relative to other Vs domains of known structure: In V\$2.3, the c" strand is hydrogen bonded to the d strand of the adjacent (outer) & shoct, whereas in other V hs the c" strand is associated with the c' strand in the same (inner) sheet (27, 29, 31, 33). A consequence of the c" strand switch is a repositioning of CDR2 and FR3, which contribute 50% and 34%, respectively, of the total contacts to SEB. It is interesting that no bacterial or viral SAGs have been described with reactivity toward members of the mouse V/12 family (3), consistent with the unique folding topology of V\$2.3. Similarly, when mouse V\$5.2 (31) is superposed onto mouse VBS.2 (Figure 4C), the RMS difference in a-carbon positions for residues in the SEB-binding site is 2.1 Å. This difference is mainly attributable to a displacement of the e" strand in a direction opposite from that of the e" strand in the mouse V\$2.3 domain, which again results in a repositioning of CDR2 and FR3. Except for MAM, which reacts with mouse V\$5.1, no SAGs specific for members of the mouse V\$5 family have been reported (3). These results indicate that the relative position of the c" strand in V\$ domains is critical in determining their reactivity toward different microbial SAGs and suggest that  $\nabla \beta$ s reactive with SEB or SEC (mouse  $\nabla \beta$ 3, 7, 10, and 17; human  $\nabla \beta$ 3, 5, 12, 13. 14. 15. 17. and 20) probably have a \(\beta\)-strand topology in their SAG-binding sites similar to that of mouse  $V\beta 8.2$ .

The structures of the  $\beta$ -SEB and  $\beta$ -SEC complexes also explain why T cells expressing mouse V\$\beta\$.2 are stimulated by SEB, SEC1-3, and SPEA, but not by SEA, SED, SEE, TSST-1, or SPEC (3,158). When SEC3 (12) is superposed onto SEB (9), the RMS difference in  $\alpha$ -carbon positions for 11 residues in the TCR-binding site is only 0.51 Å (Figure 5A, see color plates). Although the three-dimensional structure of SPEA is not known, a sequence alignment with SEB and SEC reveals that it retains several key V\$\beta\$-contacting residues, in particular Asn60, Tyr90, and Gln210 (Figure 3A). Alanine-scanning mutagenesis has shown that these three residues are not spots for the binding of SEC3 to the 14.3.d \$\beta\$ chain (111), in agreement with the fact that SAGs having other residues at these positions display different V\$\beta\$-binding specificities. Thus, SEA, SED, SEE, SPEC, and TSST-1, which do not activate V\$\beta\$2-bearing T cells, differ from SEB at nearly all V\$\beta\$-contacting positions, in particular 90 and 210 (Figure 3A). Furthermore, when SEA (14) is superposed onto SEB, the RMS difference in \$\beta\$-carbon positions for residues in the TCR-binding site

The importance of these conserved interactions to complex stabilization is demonstrated by the finding that SEC3 mutants Aso23  $\rightarrow$  Ala and Gln210  $\rightarrow$  Ala bind the TCR  $\beta$  chain 70-fold less tightly than does the wild-type SAO, whereas SEC3 Asn60  $\rightarrow$  Ala binds 16-fold less tightly (111). A recognition mechanism involving a major role for maln-chain hydrogen bonds can be highly sequence independent, enabling SEB or SEC to recognize virtually any  $V\beta$  domain in which the positions of the relevant main-chain atoms are close to those of mouse  $V\beta$ 8.2; a similar binding mode has been described for peptide/MHC complexes (144, 145). A sequence alignment of mouse and human  $V\beta$  families reactive with SEB or SEC illustrates the diversity of amino acids that can be accommodated at  $V\beta$ -contacting positions (Figure 3B).

Four water molecules were found to form hydrogen bonds bridging  $V\beta$  and SEB:  $V\beta$  Ala67 O-H<sub>2</sub>O-Oe2 Glu22 SEB,  $V\beta$  Tyr50 O-H<sub>2</sub>O-O Tyr91 SEB,  $V\beta$  Tyr65 O-H<sub>2</sub>O-Nδ2 Asn178 SEB, and  $V\beta$  Lys66 N $\zeta$ -H<sub>2</sub>O-N Phe177 SEB. Bound water molecules have also been observed in the combining site of antibodies, where they act to increase complementarity in the interface with antigen (146-148).

There are no direct contacts between SEB or SEC and  $V\beta$  CDR3, which folds away from the SAG (Figure 2A, Table 1); this is consistent with the finding that bacterial and viral SAGs stimulate T cells expressing particular  $V\beta$  elements without obvious selection for  $V\beta$  CDR3 length or sequence (3-5). However, this does not rule out the possibility that, depending on its conformation,  $V\beta$  CDR3 may in certain cases modulate SAG reactivity. For example, when the  $V\beta$  domain of TCR A6 (33) is superposed onto the 14.3.d  $V\beta$  domain in the  $V\beta$ C $\beta$ -SEB structure, SEB Tyr94 is predicted to contact Leu98 of A6  $V\beta$  CDR3, located at the tip of this long protrading loop (not shown). Similar interactions may explain the observed influence of  $V\beta$  CDR3 residues on T cell reactivity toward MAM (149) and mouse retroviral Mtv-9 SAG (150). Alternatively, these SAGs may bind the TCR in different orientations than SEB. The latter possibility is supported by the finding that reactivity to the mouse retroviral SAG MIs-1 is affected by mutations at  $V\beta$  positions 19, 20, and 24, which are not part of the interface with SEB or SEC (3, 4, 151).

### Conformational Changes in the TCR β-SEB Interface

The availability of high-resolution crystal structures for uncomplexed 14.3.d  $\beta$  chain (27) and SEB (9) permits an assessment of whether any conformational changes occur in  $V\beta$  or the SAG upon complex formation. The free and complexed  $V\beta$  domains superpose with a root-mean-square (RMS) difference of 0.33 Å for all  $\alpha$ -carbon atoms. Likewise, the unbound and bound SEB molecules superpose with a RMS difference of 0.54 Å. Thus, there are no major rearrangements in the polypeptide backbones of  $V\beta$  or SEB associated with

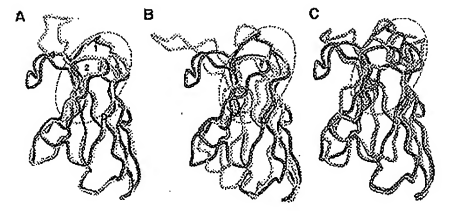


Figure 4 Comparison of  $V\beta$  structures in the region of the SAG-hinding site. (A) Mouse  $V\beta$ 8.2 (red) (27) superposed onto human  $V\beta$ 12.3 (blue) (33). The CDR loops are numbered 1, 2 and 3; HV4 in labeled 4, (B) Mouse  $V\beta$ 8.2 (red) superposed onto mouse  $V\beta$ 8.2 (red) superposed onto mouse  $V\beta$ 6.2 (graen) (31). The SEB-hinding site of mouse  $V\beta$ 8.2 is circled in each panel. The  $c^{\prime\prime\prime}$  and d strands are labeled.

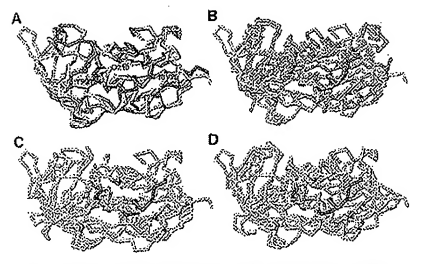


Figure 3. Structural comparison of SBB with other bacterial SAGs. SEB is oriented with its TCR-hinding site directly facing the reader. (A) a-curbon diagram of SEB (vollow) (V) superposed onto SEC3 (dark blue) (13). (B) SEB (vollow) superposed onto SEA (green) (14). (C) SIB (vollow) superposed onto TSST-1 (plnk) (1911). (D) SEB (vollow) superposed onto SPEC (light blue) (16). Regions of SEB in connect with VB in the 14.3.d \$-SEB complex we red; key connect residues are labeled.

is 2.8 Å (Figure 5B). As shown in Figures 5C and D, the putative TCR-binding sites of TSST-1 (10, 11) and SPEC (16) are markedly different from that of SEB; this can account for the finding that the VB specificities of TSST-1 and SPEC do not overlap with those of SEB or SEC (3).

### COMPARISON OF TCR-SAG-PEPTIDE/MHC AND TCR-PEPTIDEMHC COMPLEXES

Although the three-dimensional structure of a TCR-SAG-peptide/MHC complex has not been determined, a model of this complex may be readily constructed by least-squares superposition of (a) the 14.3.d VBCB-SEB complex, (b) the SEB-peptide/HLA-DR1 complex (122), and (c) the 2C TCR or heterodimer (29), which uses the same VB element (mouse VB8.2) as does 14.3.d TCR (Figure 64, see color). The accuracy of this model, in which the SAG is seen to bridge the APC and the T cell, depends on the assumption that there are no major conformational changes in any of the individual componeats upon complex formation; such changes are unlikely, given that none are observed in the TCR \(\textit{\beta}\)-SEB or SEB-peptide/DR1 complexes (37, 122). This model may be compared with the structure of the 2C TCR complexed with peptide/MHC class I (34) (Figures 6B and C). Assuming that TCRs bind MHC class I and class II molecules in similar orientations, as recently argued on the basis of structural considerations (33), it is apparent that the binding of peptide/MHC to TCR in the TCR-peptide/MHC complex is different from that in the TCR-SEB-peptide/MHC complex and that there is only partial overlap between the binding sites on the TCR for SEB and for peptide/MFIC. In the TCR-peptide/MHC complex (Figure 6B), the peptide antigen, as well as both the oil and o2 helices of the class I molecule, simultaneously engage the TCR combining site. By contrast, in the model of the TCR-SEB-peptide/MHC complex (Figure 64), the peptide is effectively removed from the TCR combining site, and there are no direct contacts between the TCR  $\beta$  chain and the MHC class II  $\alpha$ 1 or  $\beta$ 1 (which corresponds to  $\alpha$ 2 in class I) helices. However, as discussed below, the MHC BI helix is predicted to interact with the TCR Vox domain. In addition, the rotational orientation of TCR and MHC molecules in the TCR-SEB-peptide/MHC complex is different from that in the TCR-peptide/MHC complex (compare Figures 61 and C, respectively, in which the TCRs are shown in the same orientation): The MHC class If molecule in Figure 6A must be rotated approximately 40° counterclockwise around a vertical axis to align it with the MHC class I molecule in Figure 6C. Therefore, even though the TCR engages peptide/MHC differently in the two types of complexes, the end result—highly efficient T cell activation—is similar. This implies that the specific geometry of TCR engagement by peptide/MHC

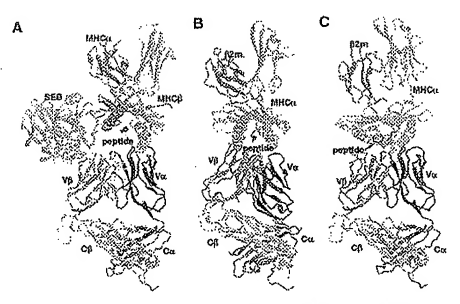


Figure 6 Comparison of TCR-SAG-peptide/MHC and TCR-peptide/MHC complexes. (A) Model of the TCR-SEB-peptide/MHC class II complex constituted by least-sequence superposition of: 1) the 14.3.d VBCB-SEB complex (37), 2) the SEB-peptide/HLA-DRI complex (122), and 3) the 2C TCR of heterodimer (29). (B) Structure of the 2C TCR-peptide/MHC class I complex (34). The complex is oriented such that the MHC molecule is approximately aligned with that in panel A. (C) Another view of the 2C TCR-peptide/MHC class I complex. The complex is oriented tuch that the TCRs in panels A and C are aligned.

may be less critical than are other factors, such as the affinity and kinetics of the binding reaction, in triggering T cells.

By acting as a wedge between the TCR  $\beta$  chain and the MHC class II  $\alpha$  chain, the SAG is able to circumvent the normal mechanism for T cell triggering by specific peptide/MHC complexes. The result is polyclonal activation of whole populations of T cells expressing particular  $V\beta$  elements, largely irrespective of the peptide/MHC specificity of the corresponding TCRs. The absence of direct contacts between peptide and TCR in the model of the TCR-SEB-peptide/MHC class II complex in Figure 6A can explain the finding that formation of a TCR-SEB-DRI complex was not affected by the presence of several different DRI-bound peptides (109). However, depending on the particular peptide bound by the MHC class II molecule, and on the conformations of the  $V\alpha$  and  $V\beta$  CDR3 loops, the peptide may in certain cases make a small number of contacts with the TCR and so modulate SAG activity.

It must be emphasized that other SAGs may bind differently to TCR and/or MHC class II than does SEB or SEC, thereby affecting the geometry of the TCR-SAG-peptide/MHC complex. In the case of TSST-1 bound to HLA-DR1 (Figure 1B), the SAG reaches across the antigen-binding groove, such that peptide-SAG and SAG-MHC  $\beta$  chain interactions may also contribute to complex stabilization (124). Furthermore, in contrast to SEB and SEC, the mode of binding of TSST-1 to DR1 would probably also preclude direct TCR-MHC interactions. This may represent an extreme example of the model in Figure 64 and illustrates how variations in the structure and positioning of the SAG wedge provide a means for different SAGs to modulate the degree of TCR-MHC interactions in the TCR-SAG-peptide/MHC complex.

## Role of the TCR & Chain in Stabilization of the TCR-SAG-Peptide/MHC Complex

There is increasing evidence that the TCR  $\alpha$  chain may, in certain cases, play a role in stabilizing the TCR-SAG-peptide/MHC complex and thereby influence T cell reactivity to bacterial or viral SAGs (5, 142, 159). For example, it was shown that  $V\alpha 4$  is expressed by  $V\beta 6^+$  T cell hybridomas that react with SEB but not by  $V\beta 6^+$  hybridomas that do not respond to this SAG (160). Transfection experiments demonstrated that the  $V\alpha 4$   $\alpha$  chain transferred SEB responsiveness regardless of whether the  $V\beta 6$   $\beta$  chain was derived from a responsive or nonresponsive hybridoma (161). These effects of the TCR  $\alpha$  chain on T cell activation by SAGs may be mediated through an interaction between  $V\alpha$  and the MHC class II  $\beta$  chain (5, 142, 161, 162). Thus, mutations at position 77 of the 1-E<sup>k</sup>  $\beta$  chain (141) and at positions 77 and 81 of HLA-DRI  $\beta$  chain (140) were found to greatly reduce the T cell response to SEB without affecting binding of the SAG to MHC class II, which suggests contacts to the TCR. In addition,

the affinity of SEB for a soluble human TCR was observed to be significantly enhanced by the addition of soluble HLA-DR1 (109). These results may be understood in terms of the model of the TCR-SEB-peptide/MMC complex in Figure 6A, in which the V $\alpha$  domain of the 2C TCR is predicted to interact with the MHC  $\beta$ 1 helix. A close-up of putative contacts (<4 Å) between V $\alpha$  CDR2 residues Ser51, Gly52, and Asp53 and the class II  $\beta$  chain residues Asp76 and Thr77 is shown in Figure 7A (see color plates). Thus, the overall stability of the TCR-SAG-peptide/MHC complex is probably determined by the combined strengths of three separate sets of interactions: TCR  $\beta$  chain-SAG, SAG-MHC  $\alpha$  chain, and MHC  $\beta$  chain-TCR  $\alpha$  chain.

The preferential expression of certain Varegions among SAG-reactive T cells has been interpreted as evidence that these particular Vox interact with MHC more favorably than do other Vas during SAG-mediated T cell activation (5, 142, 159-161). The availability of crystal structures for several aß TCR heterodimers (29-31, 33) allows an examination of the possible effects of different TCR  $\alpha$  chains on  $V\alpha$ -MHC interactions in the TCR-SAG-peptide/MHC complex. By superposing the V\$ domain of the 14.3.d \$-SEB complex onto the VB domain of TCR A6 (33), N15 (31), or KB5-C20 (30), as described above for the 2C TCR, it is apparent that the extent of interaction between the  $V\alpha$ and MHC B1 domains in the TCR-SEB-peptide/MHC complex depends mostly on the relative orientation of Ver and VB domains in each TCR heterodimer. Because the variability in the geometry of Va/VB association among these TCRs is considerable (2, 31), large differences are observed in the extent of Vα-MHC β1 interactions. These are illustrated in Figure 7. For the 2C TCR, as discussed above, Va CDR2 Ser51, Gly52, and Asp53 are predicted to contact MHC 81 Asp76 and Thr77 (Figure 7A). For the A6 TCR, Va CDR1 Gln30, and CDR2 Tyr50, Scr51 and Asn52 contact MHC \$1 Glu69, Ala73, Asp76, Thr77, and His81 (Figure 78), For the N15 TCR, Va CDR1 Lcu29 and CDR2 Thr51 contact MHC \$1 Ala73, Thr 77, and His\$1 (Figure 7C). For TCR KB-C50, no contacts are predicted because of the particular geometry of  $\nabla \alpha / \nabla \beta$  association of this TCR (not shown). In all cases where contacts between Vo and the MHC molecule are expected to occur, however, these involve Va CDR2 and residues on the MHC fil helix pointing away from the peptide-binding groove. Thus, depending on the geometry of Va/VB association and on the structure of the Ver CDR2 loop, Ver-MHC class II interactions may (a) contribute to stabilizing the TCR-SEB-peptide/MHC complex and thus increase reactivity toward the SAG. (b) have no net effect on complex stability and not affect reactivity, or (c) destabilize the complex through unfavorable contacts and thereby decrease reactivity. In this way, the TCR or chain may modulate the level of activation by SEB of T cells expressing the same V/ but different Vas.

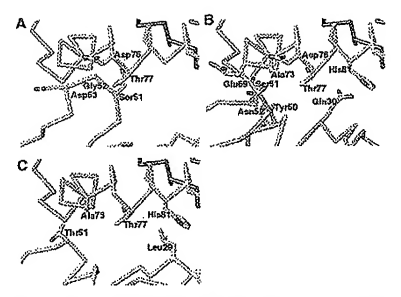


Figure 7 Differences in Va-MHC \$\textit{\textit{e}}\$ chain contacts as a result of differences in Va/V\$ orientation and Va CDR sequences. (A) Close-up of putative contacts between the MHC class II \$\textit{\textit{\textit{e}}}\$ chain Overlaw) and Va of the 2C TCR (blue) in the model of the TCR-SEB-peptide/MHC complex in Figure 6A. The bound peptide is red. Only those MHC and Va residues predicted to form direct contacts are labeled. In the upper right hand corner is a portion of the MHC class II \$\textit{\textit{e}}\$ chain and Va of the A6 TCR (arrays). The model of the TCR-SEB-peptide/MHC complex was constructed in the same way as in Figure 7A, except rating TCR A6 (33) intend of TCR 2C (29). The V\$\textit{\textit{e}}\$ domain of TCR A6 was superposed onto 14.3 d V\$\textit{\textit{e}}\$ by overlapping structurally equivalent FR residues. (C) Contacts between the MHC class II \$\textit{e}\$ chain and Va of the N15 TCR (plats). The V\$\textit{e}\$ domain of TCR N15 (31) was superposed and 16.1.3 V\$\textit{e}\$ by overlapping structurally equivalent FR residues.

# POTENTIAL THERAPEUTIC APPLICATIONS OF SUPERANTIGENS

Because bacterial SAGs are such extremely potent activators of the immune system, efforts are currently underway in a number of laboratories to engineer them for therapeutic applications (163). Knowledge of the three-dimesional structure of SAGs, and of their binding sites for TCR and MHC, may be used to design variants with altered binding properties toward these ligands, resulting in desired biological effects. The potential applications of SAG derivatives include cancer immunotherapy and the treatment of infectious and autoimmune diseases.

The recruitment of antigen-specific cylotoxic T lymphocytes (CTLs) is a major goal for the immunotherapy of malignant tumors. However, the frequency of tumor-specific CTLs is generally too low to interfere with progressive tumor growth. An attractive approach for immunotherapy is to use amibodics specific for tumor-associated antigens to target large numbers of T cells to the tumor. Taking advantage of the ability of SAGs to activate large populations of T cells, chemical conjugates of SEA and the colon carcinomareacting monoclonal antibodies (mAbs) C215 or C242 were shown to mediate T cell-dependent destruction of colon carcinoma cells lacking MHC class II molecules (164). The SEA-mAb-mediated cytotoxicity was MHC class II independent and did not require antigen-specific effector CTLs. In subsequent work, a recombinant fusion protein of SEA and the Fab region of the C215 mAb was found to efficiently target T cells to lyse C215+ MHC class II-negative human colon carcinoma cells (165). Treatment of mice carrying B16 melanoma cells expressing transfected C215 antigen resulted in 85-99% inhibition of tumor growth and allowed long-term survival. In similar experiments, SEA bound to specific anti-carcinoms cell or anti-ganglioside GD2 mAbs displayed T cell-mediated cytotoxicity toward MHC class II-negative lymphatic leukemia cell lines or neuroblastoma cells, respectively (166, 167). The demonstration of a Zn24-dependent MHC class II binding site with high effinity in the large domain of SEA (14, 128, 129), which is distinct from the low-affinity SEB-like binding site in the small domain, prompted the introduction of a point mutation (Asp227 -> Ala) in the high-affinity site in order to lower the systemic toxicity of Fab-SBA conjugates (168). Thus, after treatment with Fab-SBA Asp227 -> Ala, a 100- to 1000-fold reduction in scrum levels of IL-6 and TNF was observed in mice compared with the wild-type conjugate, without affecting anti-tumor activity. These results suggest that bacterial SAGs can be converted into tolerable immunotoxins for cancer therapy (169).

Inactivated forms of SAGs might be useful as vaccines to protect against staphylococcal or streptococcal toxic shock. Mutants of SEB that do not bind TCR do not induce T cell proliferation and therefore do not cause toxic shock (170). Animals immunized with mutants of SEA with attenuated binding to TCR or MHC class II developed high titers of anti-SEA antibodies and were fully protected against challenge with the wild-type toxin (171). Formalinized SEB toxoid-containing microspheres have been tested for efficacy in rhesus monkeys as a vaccine candidate for respiratory toxicosis and toxic shock (172). Protective immunity correlated with antibody levels in both the circulation and the respiratory tract. Similar results were obtained with intranasal or intramuscular immunization by using meningococcal outer-membrane protein proteasome-SEB toxoid. The proteasome-SEB toxoid vaccine was efficacious in protecting 100% of monkeys against severe symptomatology and death from acrosolized-SEB intoxication (173).

T cell activation by SAGs is generally followed by the disappearance or inactivation of the responding T cells, resulting in clonal deletion of cells bearing specific VH elements (3, 4). Chronic exposure to low concentrations of SAGs permits clonal deletion to occur directly, without the cells first passing through a state of hyperreactivity (174, 175). In mice with EAE or lupus nephritis, both of which serve as models for human autoimmune diseases, treatment with SEB resulted in a reduction in symptoms or in a cure of the disease (176-179). In both models, autoiromunity is known to be mediated by self-reactive T cells expressing a single VB element (mouse VBS), such that the effect of the SAG most likely results from the specific elimination of pathogenic T cells bearing that particular VB. These experiments suggest that SAGs could potentially be used for the prevention of autoimmune disease by selectively climinating specific T cell populations. In cases where T cells from several VB families might be involved in the disease process, SAGs engineered to recognize these Viss could be used for therapy. Knowledge of the three-dimensional structure of TCR-SAG complexes should facilitate the design of SAGs with predefined VB specificities.

A number of concerns must be addressed, however, before SAGs can be used as therapeutic agents (180). Administration of SAGs may lead to the release of dangerous levels of cytokines such as TNF, resulting in toxic shock. In addition, inadvertent stimulation of autoreactive T cells could trigger autoimmune disease. It appears likely, however, that the systemic toxicity of SAGs can be dissociated from the superantigenic effects of these molecules through structure-based genetic engineering, as described above for SEA-Fab conjugates (168) and SEA vaccines (171). Similarly, mutations of TSST-1 have been described that alter either lethality or superantigenicity, without significantly affecting the other property (181–183).

#### **FUTURE DIRECTIONS**

Although the X-ray crystallographic studies described above represent important advances in our understanding of SAG interactions with TCR and MHC, the apparent diversity of these interactions clearly illustrates the need for further work in this area. For example, TSST-1 binds differently to MHC class If molecules than does SEB (Figure 1) (122-124), and its interaction with the TCR presumably differs as well. Unlike SEB and TSST-1, which have only one class II-binding site, mutagenesis and biochemical experiments indicate that SEA has two such sites (127-129). However, because the crystal structure of an SEA-MHC class II complex has not been determined, the precise locations of these sites are unknown. SED and SPEC crystallize as homodimers (15-16). which may facilitate TCR oligomerization and T cell triggering. However, X-ray crystallographic studies are required to clucidate the geometry of the putative SED, TCR2, SED, MHC2, SPEC, TCR2, and SPEC, MHC2 tetramers. Our knowledge of SAG structure is currently limited to the staphylococcal and streptococcal pyrogenic toxins (9-16) and to a staphylococcal exfoliative toxin (17, 18). No structural information is available for SAGs produced by mycoplasma, such as MAM (19), or by viruses (22-26), including the muchstudied MMTV SAGs. Human endogenous retroviral SAGs that are associated with autoimmune diseases (26) may become a focus of future attention. Finally, the structures of entire TCR-SAG-MHC complexes must be determined in order to define experimentally the putative Va-MHC interactions discussed in this review.

X-ray crystallographic studies of TCR-SAG, SAG-MHC, and TCR-SAG-MHC complexes will open the way for the design of SAG variants with altered binding properties for TCR and MHC for use as tools in dissecting structureactivity relationships in this system. The relative contributions of TCR-SAG and SAG-MHC interactions to T cell stimulation can be defined by engineering panels of mutant SAGs with both higher and lower affinities for TCR and MHC than the wild-type toxins have. It has been shown that SAGs mimic the interaction of peptide/MHC complexes with the TCR in terms of affinities and kinetics (109-112). It remains to be established, however, whether there is an optimum affinity for T cell activation by SAGs (or by peptide/MHC), as predicted by the serial triggering (118, 119) and kinetic proofreading (120) models of T cell activation, such that SAGs with either higher or lower affinities than this optimun value exhibit decreased ability to stimulate T cells. Alternatively, SAGs with progressively higher affinities for the TCR relative to the wild type may stimulate T cells increasingly well, until some plateau of maximum stimulation is attained. Mutants with altered affinities for the TCR can be used to distinguish between these possibilities and thereby define the affinity and kinetic parameters governing T cell activation by SAGs. The role, if any, of cooperative interactions involving the TCR  $\alpha$  chain in stabilizing the TCR-SAG-MHC complex can be similarly addressed. By thus combining X-ray crystallography with mutagenesis and binding studies, a comprehensive understanding of the physical basis of T cell activation by microbial SAGs should emerge.

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#### Literature Cited

 Bentley GA, Mariuzza RA. 1995. The structure of the T cell entirest receptor. Anna. Rest Immunol. 14:563-90
 Wilson IA, Gortin KC. 1997. T-tell recep-

 Wilson IA, Gortin KC, 1997. T-nell recepter structure and TCR complexes. Curr Optn. Struct. Biol. 7:839-48

3. Kotzin BL, Letrng DYM, Knippler J, Macrack P. 1993. Supermitigens and their potential role in human disease. Adv. Internal., 54:99-166

 Schieter MT, Ispatowicz L, Winslow GM, Kappler JW, Marnek P. 1993. Superandgens: bacterial and viral proteins that manipulate the lumence system. Aumt. Res Cell. Biol. 9:101-128

 Webb SR, Geneoigno NRI. 1994. T-cell netivation by superantigens. Curr. Opin. Immunol. 6:467-475

 Bohach GA, Fast DJ, Nelson RD, Schlievert PM. 1990. Staphyloooccal and stroptococcal pyrogenic toxins involved in toxic shock syndrome and related illnesses. Crit. Res. Microbiol. 17:251-272.

 Betley MJ, Berst DW, Regates LB. 1992. Staphylococcal enterocoxias, todo shock syndrome taxin and streptococcal pyrogenic explanine: a comparative study of their molecular biology. Chem. Immunol. 521, 126

 Bohach GA. 1997. Stephylotoccal enteroloxins B and C. In Supercritiques: Malecular Biology, Immunology and Reference, to Human Discota, ed. DYM Leung, BT Huber, PM Schlieven, Marcel Dekker Inc. NY, pp. 167-198 Swemingthan S, Fatey W, Pletcher J, Sax

 Sweminethan S, Fairny W, Pletcher J, Sax M. 1992. Crystal structure of stephylococcal enterotoxin B, a superantigen. Natture 359:801-806

10. Prisad GS, Eathart CA, Murray DL, Novick RP, Schlievert PM, Oblendorf DH. 1993. Structure of taxic shock syndrome toxin-1. Biachemistry 32:13761-13766.

 Acharya KR, Possalacqua EF, Jones EY, Harlos K, Stuart DI, Brehm RD. Trance HS. 1994. Structural basis of superantigen action inferred from crystal structure of toxic-shock syndrome toxin-1. Nature 347:94-97

 Hollmann ML, Jablonski LM, Crum KK, Hackell SP, Chi Y-I, Stauffacher CV, Stevens DL, Bobach GA. 1994. Predictions of T cell receptor and major histocompatibility complex binding sites on staphytococcal enterotoxin C3. Infact. Imman. 62:3396–3407

 Papareorgiou A, Acharya KR, Shapiro R, Passolacqua EF, Brehm RD, Tranter HS. 1995. Crystol structure of the superantgen enterotoxin C2 from Stapylococous aureus reveals a vine-binding site. Structure 3:760-779

 Schad EM, Zaitsev J, Zaitsev VN, Dohlsten M, Kolland T, Schlievert PM, Ohlendorf DH, Svensson LA, 1995.

- Crysul structure of the superantigen staphylococcol enterotoxin EMBO J. 14:3292-3301
- Sundstrom M. Abrahansen L. Antoneson P. Mehindate K. Mourad W. Dobisten M. 1996. The crystal structure of staphylocoreal enterotoxin type D reveals Zn2-mediated homodimerization. EMBO J. 15:6832-6840
- Roussell A, Anderson BF, Baker HM, France ID, Baker EN, 1997, Crystal strucane of the anertocoreal supercratizen SPE-C: dimerization and sinc-binding suggest a novel mode of interestion with MHC class II molecules. Nature Struck Hiol. 4:635-649

With GM, Eurhot CA, Rago IV, Kim MH, Behach GA, Schlievert PM, Ohlendart DH, 1997, The structure of the superantigen exfolintive toxin A suggests a novel reguintion us a serine protesse. Blochem-tiny 36:1559-1566

Cavarelli J. Prevost G. Bourgurt W. Moulinier L. Chevrier B. Delagoutte B. Bluves A. Mourey L. Rifan S. Piemont Y. Moras D. 1997. The structure of Staphylococcus anneus epidermolytic toxin A. an atypic serine protesse, at 1.7 A resolution. Structure 5:813-824

19. Cole BC, Griffiths MM. 1993. Triggering

and expectation of autoimmune withings by the Mycoplasma arthritidis supermiti-gea MAM, Arthritis Rheumat. \$6:994-

20. Ho Y, Abe J, Yoshino K, Takeda T, Kushaka T, 1995. Sequence analysis of the gene for a novel superandem pro-duced by Yerziula pseudotuberculosis and expression of the recombinent protein. J. Inimunol. 154:5896-5906

Miyoshi-Akiyama I, Abe A, Kato H, Koushina K, Narimatsu H, Uchiyama T, 1995. DNA sequencing of the gene encoding a bacterial tunerantigen, Yersinia psyudotuborculoxia-derived mitogen (YPM), and characterization of the gene product, closed YPM. J. Institutol. 154:5228-5234

Acha-Orben H. MacDonald HR. 1995. Superentigens of mouse mammary turnor virus. Annu. Rox Immunol. 19:459-486

Astoni E, Lofage M, Lafon M. 1996. Rabics supermotion as a VA T-dependent adjuvant. J. Sop. Med. 183:1623-1631
Debroseu D, Ursea B, Pope M, Asch AS, Poanen DM. 1995. Enhanced HV-1

replication in VBI 2 T cells due to human cylomegalovirus in mosocytes: evidence for a putative herpesvirus superantiges. Cell 82:753-763

- Sutkooreki N, Palkama T, Ciurli C, Sekal R-P, Thoricy-Lawson DA, Huber B. 1996. An Epstein-Barr virus-casoristed superantigea. J. Exp. Med. 184:971-980
   Conrad B, Weissmahr RN, Boul J, Arcari R, Schiphach J, Mach B. 1997. A hu-man and control of control superantisen.
- man endogenous retroviral superantigen as candidate outeinmune gene in type I diabeten. Cell 90:303-313

Bentley GA, Houlet G, Karjalainea K, Marjuzza RA. 1995. Crystal structure of the B chain of a T cell antigen receptor.

Science 267:1984-1987

 Fields BA, Ober B, Malchiedi EL, Lebe-deva MJ, Braden B, Ysern X, Kim J-K, Shao X, Ward EB, Marinaza RA, 1995. Crystal structure of the Var domain of a T cell unugen receptor. Science 270: 1821–1824

Garcia KC, Degano M, Stanlicid RL, Brummak A, Inckson MR, Peterson PA, Teyron L, Wilson IA. 1995, An aft T cell cocapios structure at 2.5 Å and its orientation in the TCR-MHC complex. Science 274:209-219

Plousset D. Mazza G. Gregoire C. Pirot C. Molisset B. Fonteeiths-Comps JC. 1997. The tirre-dimensional structure of a T-cell energen receptor VaVB heterodimer reveals a novel arrangement of the VI domain. EMBO J. 16:4205-4216

- the Vs domain. EMBOJ. 16:4205-4216
  Wang I, Lim K, Smolyar A, Teng M, Liu
  I, Tec AGD, Liu I, Fluetey RE, Chishi
  Y, Thomson CTT, Sweel RM, Nathenton SG, Chang H, Sachentini JC, Roinherz EL. 1998. Atomic structure of an

  aft T cell receptor (TCR) beterodimer
  in complet with an auti-TCR Fob fington the structure of the property of the structure. ment derived from a mitogenic ambody. EMBO J. 17:10-26
- Li H, Lebodeva ML, Llera AS, Fields BA. Brenner MA, Mariuzza RA. 1998. Struc-ture of the VS domain of a human yo T-cell entigen receptor. Nature 391:502-506
- Garboczi D.N., Ghosh P. Uz U., Fan QR., Blddison WE, Wiley DC. 1996. Sinicitic of the complex between human T-cell receptor, viril peptide and HLA-A2. Na-
- Garcia KC, Degano M, Pease LR, Huang M, Peterson PA, Teyton L, Wilson IA. 1998. Surcentral basis of plasticity in T cell receptor recognition of a self peptido-MilC antigen. Science 279: 1166-1172
- Ding Y-H, Smith KJ. Garboczi DN. Utz. U, Hiddison WE, Wiley DC. 1998. Two human T cell receptors bind in a simi-lar diagonal mode to the HLA-A2/Inx

peptide complex using different TCR amino soids. Immunity 8:403-411
Fields BA, Matchiodi EL, Li H-M, Ysem X, Staussacher CV, Schlievert PM, Karjeloinen K, Marineza RA, 1996, Crystal structure of the B chain of a Y-oell receptor complexed with a superantigen. Na-

nora 384:188-192 37. Li H, Liera A. Tsochiya D, Leder L, Ysern X, Schlieven PM, Karjalainen K, Meriuzzn RA. 1998. Three-dimensional struc-

uzza RA. 1998. Three-dimensional structure of the complex between a T cell reaceptor R chain and the supersettion stephylococeal enterotoxia B. Immunity MacDoneld HR. Lees RK. Baschieri S. Herrmann T. Lussow AR. 1993. Peripheral T-cell reactivity to bacterial superantigens in vivo: the response/anergy paradox. Immunol. Rev. 133:105-117 Woodhand DL. Wea R. Blackman MA. 1997. Why do supersatigeas care about peptides? Immunol. Today 18:18-22 Kappler JW. Rochan N. Martack P. 1987. T cell tolerance by clonal elimination in

Appier Jw, Rochm N, Marrack F. 1987.
 T cell tolerance by cloud elimination in the thymus. Call 49:273–280

 ManDonald HR, Schweider R, Loce RK, Howe RC, Acha-Orben H, Festenstein H, Zinkernagel RM, Hengarther M. 1988.
 Feell receptor VB use predicts reactivity and tolerance to Miss-encoded untigens.

end toterance to Misa-encoded amogens, Noture 332:40-45

42. Pailen AM, Bill J, Kusho RT, Marrack P, Kappler JW. 1991. Analysis of the interaction site for the self superanigen Mis-1A on T cell receptor VB. J. Exp. Med. 173:1183-1192

43. White J, Hennan A, Pullen AM, Kubo R, Kuppler JW, Marrack P. 1989. The VB-tyrofic superangen standardocodeal ca-

- specific superantigen staphylococcal enterotoxin B; slimulation of matuo T cells and cloud deletion in recordal mice. Cell 56:27-35
- 50:27-35
  Rellahan BL, Jones LA, Kralsback AM, Fry AM, Malis LA. 1990. In who induction of anergy in peripheral V/18<sup>+</sup> T cells by staphyloconcal enterotoxin B. J. Exp. 14td, 172:1091-1100
- O'Hehir RE, Lamb JR. 1990. Industion of specific closel spergy in human T tym-phocytes by Staphylococcus arrens en-terotoxins, Proc. Natl. Acad. Sci. USA 7:8884-8888
- Kawabe Y, Othi A. 1991. Programmed coll death and extrathyonic reduction of VAS\* CD4\* T cells in mice tolerant to Staphylococcus aureus enterotoxia B.
- Notice 349:245-748
  Golovica TV, Chervonsky A. Dudley IP,
  Ross SR. 1992. Transgene mouse man-mary tumor virus superantigen expression provents viral infection. Cell 69:637-645

Reid W. Wannders G. Shabov AN, Scarpelline L. Acha-Orbea H. MacDon-ald HR. 1995. Superantigen-fedured im-mune stimulation amplifies mease man-nione.

mary tumor virus infection and allows virus tumoriusino. Cell 74:529-540 lgaatowicz L. Kappier I. Marreck P. 1992. The effects of chronic infection with

Mod 175:917-923
Golovkins TV, Dudley IP, Jaffe A, Ross SR. 1995. Mouse mammay mmor viruses with finactional supprantigen genes are sciected during in vivo infos-tion, Proc. Natl. Acad. Sci. USA 92:4828-4892

Wrona TJ, Lorano M, Binhazim AA, Dudley JP, 1998. Mutetional and functional analysis of the C-terminal region of the C3H meass mammary hazor vices appearation. J. Vind. 72:4746-4755

Deresiewicz RL. 1997. Staphylococcal toxic shock syndroma. In Superantigens: Malaculor Biology, Immunology and Ral-evence to Human Disease, ed. DYM Le-

evance to Human Discore, ed. DYM Leung, BT Huber, PM Schileven, Marcel Dekker Inc. NY, pp. 435-479
Stevens DL. 1997. Streptococcal taxic shock syndrome. In Superantigens: Molocular Biology, Immunology and Relevance to Human Discore, ed. DYM Leung, BT Huber, PM Schileven, Marcel Dekker Inc. NY, pp. 481-501
Rago IV, Schilevan PM. 1998. Mechanisms of puthogenesis of simply/ococcal and streptococcal superantigens. Car. Top. Microbiol. Immunol. 225:81-97
Alber G, Hemmer DK, Fleiseler B. 1990. Relationship between enterotoxio-

- 1990. Relationship between enterotoxio-and T lymphocyte-stimulating activity of stephylospooni enterotoxin. Br. J. Im-
- of supplysopeon entertown. Dr. 3, Internation 144:4501–4506
  Harris TO, Grossman D, Koppler IV, Marzack P, Rich RR, Heltey MJ. 1993.
  Lack of complete correlation between emetic and T cell sliminary polytics of
- emetic and text standards acceptance of suphylococcal enterotoxins. Infect. Immun. 61:3175-3183

  Hovde CJ, Mart JC, Hoffmann ML, Hacken SP, Chi YI Crum KK, Stovens DL, Stauffacher CV, Bohneh GA. 1994. Investigation of the tigation of the role of the distribilde bond in the activity and sinceture of staphy-torocent enterclosin Cl. Mol. Microbiol.
- Sears CL. Kapar JB. 1996. Enteric bac-terial toxine: mechanisms of action and linkage to intestinal socretion. Microbiol. Rev. 60:167-215
- Hamed ARA, Mariack P. Kappler IV. 1997. Transcytosis of staphylococcal

superantigen toxins. J. Exp. Mod. 185:

superanuges toxics. A personal of the control of th Sci. 475:146-156

Ray CG, Palmer IP, Crossley JR, Williams RH, 1980, Coxenckie B virus antibody responses in juvenile-opset disbeles mellitus. Clin. Endocrinol. 12:375-

Oldstone MBA. 1998. Molecular mittai-cry and autoimmune disease. Cell 50: 819-830

819-820
Witcherpfenning KW, Stronninger II.
1995. Molecular mimiery in T-cell mediated autoimmutality: viral peptides activate human T cell clones spenific for
myelin basic protein. Cell 80:895-705
64. Hommer B, Fleckenstein BT, Vergelli M,
lung G, Molectand H, Martin R. 1997.
Identification of high potency microbial
and self ligands for a human autoreactive class II-restricted T cell clone. J. Exp.
Med. 185:1651-1659

Med. 185:1651-1659
65. Remo T, Acha-Oibea H. 1996. Superantigets in autoimmuna disease: still more shades of gray. Immanol. Rev. 154: 175-191

66. Bush JF. 1995. T cell receptor use in organ-specific human autommune dis-enses other than theumatoid arthritis and multiple selemis. Arm. N.Y. Acad. Sci. 756:453-459

67. Trangon U. 1987. Multiple scienosis: relevence of class 1 and class H MHC-expressing cells to leston development J. Neurotinaturol. 16:283-302 68. Olemp O, Hillert J. 1990. HLA class H

associated genetic susceptibility in multi-ple sciences: a critical evaluation. Tisma Antigens 98:1-5

Allegretts M, Nickles IA, Sniram S. Albertini RJ. 1990. T cells responsive to

Albertial RJ. 1990. T cells responsive to myelin basic protein in patients with multiple sciences. Science 247:718-721 Wacherpfenelg KW. Zhang J, Witck C, Matwi M, Modabber Y, Ota K, Hafter DA. 1994. Cloual expansion and pertistance of human T cells specific for an immunodominant myelia basic protein peptida. J. Immunol. 150:5581-5597. Zhang J, Markovic S, Lacet B, Rans J, Weiner HL, Hafter DA. 1994. Increased feequency of interieudin 2-responsive T cells specific for myelia basic protein

I cells specific in: myelin busic protein and prokolible protein in peripheral blood and cerebrospinal fluid of patients with multiple reterosin. J. Exp. Med. 179: 72. Stiniesen P. Reur J. Zhang J. 1997. Au-toimmune pathogenesis of multiple sele-rosis: role of autorecutive T lymphocytes

and new immunotherapeutic strategies. Cell. Rev. Journal, 17:33-75
Brocke S. Grar A. Pierov C. Gratam K. Gilbels C. Fathman CG, Steinman L. 1993. Induction of relapsing pural-

L. 1993. Indiction of relapsing paralysis in experimental autoimmune encephatomyelitis by becterial supermainen. Nature 365:642-644
Rocke MK, Quigley L, Cannella B, Raine C, McFarlin DE, Scott DE, 1994. Supermitigen modulation of experimental allergic encephatomyelitis: antivation or proportional entering and property.

icipic encephatonyculot: 3 divation of interpolation of interpolations outcome. J. Immunol. 152:2051-2059

Brooke S, Hausmann S, Stelmman L, Wicherpfennig KW, 1998. Microbial pepides and superanigens in the pathonomic of interpolations. genesis of autoimmune diseases of the central nervous system. Sem. Immunol. 10:57-67

 Palierd X, West SG, Lafferty IA. Clements IR, Keppler JW, Marrack P. Kotzin BL. 1991. Evidence for the effects of a superantigen in rheumateid arthritis. Spionee 253:325–329

Hagni TM, Anderson GD, Bouegee S, David CS. 1992. Restricted heterogeneity in T-cell antigen receptor VB gene usage in the lymph nodes and artificiants of mice. Proc. Natl. Acad. Sci. USA 80:1252-1252.

89:1253-1255 Osman GE, Toda M, Kanngawn O, Hood LE, 1993. Characterization of the Terlireceptor repenolise sausing colingen arabritis in mice, J. Exp. Mod. 177:387-

Chioschin G. Boissier MC. Fournier C. 1991. Thirmpy against musine collagen-induced authority with T-cell receptor

induced atthritis with 1-cent receptor VB-specific antihodies. Eur. J. Intumunol. 21:2899–2905
Moder KG, Lubra HS, Galliths M, David CS, 1993. Prevention of collagenthricis of Toell receptor VBS bearing Toells with monoclonal antibodies. Br. J. Rheumol.

monocional annovates. B. J. Relation.
32:26-30
81. Kurhar V. Stream B. 1993. The involvement of T cell receptor peptide-specific regulatory CD4. Teells in recovery from unitgen-indused autoimmune disease. J. Exp. Med. 178:989-916
82. Kurhar V. Stellrecht K. Sercarz E. 1996. Inactivation of CD4 regulatory T cells are the content in temporary in J. Exp.

results to chronic automorphity. J. Exp. Med 184:1609-1617

Kumai V, Aziz F, Sercerz E, Miller A. 1997. Regulatory T cells specific for the

same Gamework 3 region of the V18.2 chain are involved in the control of colle-Ben Il-induced estimitis and experimental nutoimmuse encepholomyclids. J. Exp. Med. 185:1725—1733

Singh B, Prange S, Javnikur AM. 1998. Projective and despueline effects of microbial infection in insulin-dependent di-aborer mellitus, Som. Immunol. 10:79-

Conred B, Weldmann E, Trucco G, Rudert WA, Behboo R, Ricordi C, Rodinguez-Rlio H, Finegold D, Trucco M. 1994. By-idence for superartigen involvement in insulin-dependent diabetes mellinu actiology. Nature 371:351-355
 Luppi P, Trucco M. 1996. Superantiscus in insulin-dependent diabetes mellinus. Springer Samin. Immunopathol. 17:333-362

87. Abe J. Koizin BL. Jujo K. Melish ME. Glode MP. Kobsain T. Leune DY. 1992. Selective expansion of T cells expressing T-cell receptor variable region VBZ and VBS in Kawasaki disease. Proc. Natl.

and VBS in Kawaraki disease. Proc. Natl. Acad. Sci. USA 89:4066-4070

88. Abe J. Koton BL., Meissner C., Melish ME, Takahashi M, Pulton D, Romagne F, Molisren B, Leung DYM. 1993. Characterization of T cell receptor changes in noute Kawasaki disease. J. Bap. Med. 127:401-406. 177:791-796

Curtis N. Zheng R, Lamb JR, Levin M. 1995. Evidence for a superantigenmediated process in Kawasaki disease. Arch. Dis. Child. 72:308-311
 Leung DY, Giorno RG, Kazemi LV, Flyan PA, Busse JB. 1995. Evidence for super-

antigen involvement in cardiovasoular in-

- singen incorrement in citalovasous ra-jury due to Kawasiki syadrome. J. Im-muol. 155:5018-5021 Leung DY, Meissner HC, Polton DR, Murray DL, Kotela BL, Schlievert PM. 1993. Toxic shock syadrome toxin-secreting Stophylococcus aureus in Kawasaki syndrome. Loucet 342:1385-1488 1388
- Pietra BA, De Innocacio J. Giambini EH. Hirsch R. 1994. TCR V bele family reper-toire and T cell activation warkers in Kangasaki distance. J. Immunol. 153:1861-
- 93. Sakagushi M, Kato H, Nishiyori A, Sagawa K, Itoli K. 1995. Characterization of CD4\* T helper cells in patients with Kawasalii disease (KD): preferential production of aunor necrosis factoralpha (TNF-alpha) by Vocta2 or Victa8 CD4\* T kelper cells. Clin. Exp. Immunol. 2023. 282. 99:276-282
- 94. Juson J. Montana E. Donald JF, Suidouan

M. Inge KL. Campbell R. 1998. Kawasaki disease and the T-cell untigen receptor. Hum. Immunol. 59:29-38

95. Inadoto J. Chapes SK. 1997. The exfolistive toxins of Staphylococcus auteus. In Superantigens: Molecular Biology, Im-munalogy and Rolevance to Human Dis-case, ed. DYM Leung, BT Huber, PM Schilevert, Marcel Deider Inc. NY, pp. 231-255

Prevost G, Rifai S, Choix ML, Plemont Y. 1991. Functional evidence that the Ser-195 residue of stophylococcal exfo-

ser-193 residue of stophyloooccal extolative toxin A is estential for biological
activity. Infect. Immun. 59:3337-3339

97. Herderson CA, Highet AS, 1985. Acute
psoriasis apsorbated with Lancefield
group C and group G entancous streptocooccil infections, Ba, J. Dermotol. 118: 559-562

98. Lewis HM, Baker BS, Bokth S, Powles AV. Gurioch II, Voldinarsson H, Fry L. 1993. Restricted T cell receptor VA gene usage in the skin of patients with guttate and chronic plaque provincis. Br. J. Duratot. 129:514-520

hatol 125151-520 Chars ICC, Smith LR, Froning KI, Schwabe BJ, Laker JA, Caralli LL, Kur-land HH, Karasek MA, Wilkinson DJ, Carlo DJ, Brostoff SW, 1994. CD3 Corlo 191, Brotton 1891, 1994. Con T cells in psoriatic lessons preferentially use Feel receptor Vbeta 3 and/or Vbeta 13.1 genes. Proc. Natl. Acad. Sel USA 91:9282-9236. Loung DYM, Travers IB, Gierno R. Nerdis DA, Skinner R. Aelion J, Køzemi LV, Kim MH, Tramble AE, Kolb M, Schliever DM, 1905. Evidence for extended.

ert PM. 1995. Evidence for stropiococcal superantigen-driven process in acute guitate psoriasis. J. Clin. luvest. 96:2106-2112

2112
101. Bochneke WH, Dressel D, Manfrus B, Zolliner TM, Wettstein A, Bohm BO, Sterry W. 1995. Teell reperiotic in chronic plaque-surge poolesis is restricted and locks enrichment of superantigen-accordated VI regions. J. Invest. Darinatol. 104:725-728 K, Hanskawa Y, Sugai M, Hashimolo K. 1998. Superantigen production by Stophylococcur au-

cen production by Stophylococcus au-

198 103. Leung DY, 1995. Atopic dermatitis: the skin as a window into the pathogenesis of change allergic diseases, J. Allergy Clin. Immunol, 96:302-318

Immunol, 90:305–318 Leyden H, Marples RR, Klittgman AM. 1974. Staphylococcus await in the le-sions of atopic detrastits. Br. J. Demalal. 90:525–530

105, Leung DY, Harbeck R, Bina P, Reisir RF. Yang E, Norris DA, Harifin IM, Sampsen HA. 1993. Presence of the antibodies to stephylococcal exotoxins on the skin of supplylosoccat exclusion on the stat of pulsens with atopic dermatitis: avidence for a new proto of altergens. J. Clin. Invest. 92:13/4-1380
Gasonigne NRJ, Ames KT. 1991. Direct binding of secreted T-cell receptor fichaln

to superantigen associated with class II major histocompatibility complex protein. Proc. Natl. Acad. Sci. USA 88:613-

107. Malmqvin M. 1993. Surface pharmon res-

Malmqvia M. 1993. Surface plasmon resonance for denotion and measurement of antigen-militody affinity and kinetics. Curr. Blol. 5:282–286. Marguliet DH, Flackin D, Khilko SN, Jelonek MT. 1996. Studying interactions involving the T-cell antigen receptor by surface plasmon resonance. Curr. Optic. Immunol. 8:262–270. Seth A, Stem LS, Ottenhoff TRM, Engel 1. Owen MJ, Lumb JR, Klausner RD, Wiley DC. 1994. Binary and ternaty complexes between T-cell receptor, class II. MHC and superantigen in vitro. Nature MHC and superantigen in view. Nature 369:324-327

110. Melchiedi EL, Eltenstein E, Fields BA,
Ohtendorf DH, Schlievert PM, Karjalainen K, Mariuzza RA, 1995. Superanilgen binding to a T cell recentor fi
chain of known three-dimensional struc-

cash of known and and state and a solution. J. Exp. Med. 182:1833-1845 Loder L. Liera A. Lavoie FM. Lobedeva Mi, Li H. Sekely R.P. Bohzeh OA, Gabr PJ, Schlieven FM, Kurjalainen K, Mar-iuzza RA. 1998. A mulational analysis

iuzza RA. 1998. A mutational analysts of the binding of suphylococcal enterowins B and C3 to the T cell receptor B choin and major histocompatibility complex class H. J. Exp. Med. 187:823-833 Khandeker SS. Brauer PP, Naylor JW, Chang H.-C. Kern P. Newcomb JR, LeClair KP, Saump MS, Benencoun BM, Kawasaki E, Braceji J, Profy AT, Jones B. 1997. Affinity and kinetics of facilitaria-tions between a week T-cell in the content and is

1997. Adminy and senerce of neutral-tions between an affire it receives proposed is supersatigen and class II-MHC/peptide ligands. Mol. Immunol. 36:493-503 van der Metwe PA, Brown MH, Davis SI, Barchay AN. 1993. Affinity and isner-ies of the interaction of the cell adhesion molecules rat CD2 and CD48. EMBO J.

12:4945-4954

om der Merwe PA, Baretay AN, Ma-son DW, Davis EA, Morgan BP, Tone M, Krishnim AKC, Ianelli C. Davis S1, 1994. Human cell-adhesion moleculo CDZ binds CDS8 (LFA-1) with a very low affinity and un extremely last disso-

ciation rate but does not bind CD48 or CD59. Blocheminary 23:10149-10160 Corr M, Slanctz AE, Boyd LF, Jelenek MT, Khilko S, Al-Rumedi BK, Kim YS, Maher SE, Bothwall AL, Margulies DH. 1994. T cell receptor-MHCchass I peptide interactions: affinity, kinetics, and specificity. Science 265:946-948 Mattai K, Boallace JJ, Steffner P, Resy PA, Davis MM. 1994. Kinetics of T cell receptor binding to peptidoff-E' complexes: correlation of the distorbinda rate with T-cell reepossiveness. Proc. Nat.

meners correlation of the differsimble fall with Teell responsiveness. Proc. Nat. Acad. Sci. USA 91:12862-12866.
Alam SM, Travers PJ, Wing JL, Nasholds W, Reciputh S, Ameson SC, Cascolgae NRJ. 1996. Toell receptor affinity and dismacyte positive selection. Nature 38:1616-620.

381:016-640 Whithith S. Muiler S. Celin M. Padovan E. Lanravecchia A. 1995. Seriet Inggering of many T-cell receptors by a few peptide-MHC nomplexes. Nature 375:148-

Viola A, Langavecchin A. 1996. T cell activation determined by T cell recognor number and tunable thresholds. Science 273:104-106

Rabinovine JD. Beccon C. Lyons DS. Drois MAI, McConnell HM. 1996. Ki-netic discrimination in T-cell activation.

Proc. Notl. Acad. Sci. USA 93:1401–1405
Prof. T. Fraser J. 1998. Superanigens:
[ust like peptides only different. J. Ergs.
Med. 187:819–821

handersky TS. Brown IH. Gorga IC. Us-ban RG, Chi YI. Stauffacher C, Stro-minger IL. Wiley DC. 1994. Three-dimensional structure of a human class

dimensional structure of a human cines this monocompanibility molecule complexed with superantigen. Nature 368:711-718 Dessen A, Lawrence CM, Cupo S, Zaller DM, Wiley DC, 1997. X-ray crystal structure of HLA-DR4 (DRA\*0101, DRB1\*0401) complexed with a peptide from human collagen II. Immunity 7:473-821

124. Kim J. Urben RG, Strominger JL, Wiley DC. 1994. Toxic shock syndrome toxin-1 complexed with a class II mejor birto-compatibility molecule HLA-DR1. Science 266:1870-1874

ence 2bb:1874-1874
Mollick JA, Chistigampala M, Cook RG, Rich RR, 1991, Stephylococcal exotoxin sctivation of T-cells. Role of exotoxin-MHC clase II binding affinity and class II isotyps. J. Innumol. 146:463-468
Wen R, Cole GA, Surman S, Blackman MA, Woodland DL. 1996, Major historymanical interactions.

compatibility complex class II-associated peptides control the presentation of

bacterial superantigens to T cells. I. Exp.
Med. 183:1063-1092

127. Kordno H, Parker D, White I, Marrack P,
Kappler J. 1995. Multiple binding sites for
bacterial superantigens on soluble class B
molecules. Junuarity 3:187-196

128. Hudson KR, Tiedemann RE, Urbon RG,
Love SC, Strominger H., Fraser JD,
1995. Staphylocopeal enteriourin A bas
but concernity blading sites on major two cooperative blading sites on major

histocompatibility complex class II. J. Epp. Med. 182:711-720

129. Abrahmsen L. Dobisten M. Sagren S. Bjork P. Jonsson E. Kalland T. 1995. Chameterization of two distinct MHC class II binding sites in the superantigon staphylocopeal enterotoxin A. EMBO J. 14:2978-3986

14:2978-2986
130. Tiedemain RE, Urban RJ, Strominger PL, Frazer JD. 1995. Isolation of HLA-DR L(staphylonoceal enterotoxin A)2 vimers in solution. Proc. Natl. Acad. Sci. USA 92:12150-12159
131. Mehindate K. Thibodeau I, Dohlsten M, Kalland T, Sekaly R-P, Mourad W. 1995. Cross-linking of major histocompatibility complex class II molocules by staphylococal enterotoxin A is a scoulcursent

ity complex class II molecules by simplylococcal enterotoxin. A is a regularment
for inflammatory cylokine sene expression. J. Exp. Med. 182:1573-1577

132. Tredemain RE, Fraser ID. 1996. Crosslinking of MHC class II molecules by
tupphylococcal enterotoxin. A is essential
for antigen-presenting cell and T cell activation. J. Immunol. 157:3958-3966

133. Winslow GM, Marrack P, Kappler IW,
1994. Processing and major histocompstholity complex binding of the MTV7 superontigen. Immunol. 1:23-33

134. Mottershead DG, Hau P-N, Urban RG,
Strominger IL, Buber BT. 1995. Direct
binding of the Miv7 superantigen (Misjournelly 2:149-154

nty 2:149-154
135. Held W. Waardem GA, MacDonald HR, Acha-Orbea H. 1994. MHC class II hierarchy of supranningen presentation predicts efficiency of infection with mouse

manuscript tumor virus. Int. Immunol. 6:1403-1407

136. Pucillo C, Cepeda R, Hoder RJ. 1993. Expression of a MHC class II transgene determines superantigenicity and inscepti-

bility to mouse mammery temor virus in-fection. J. Exp. Med. 178;1441-1445 137. Thibodeau I, Labrocque N, Denis F, Hu-ber BT, Sekaly R-P, 1994. Binding sites for bacterial and endogenous reduviral supergonigens can be dissociated on mojor histocompatibility complex class II molecules. J. Exp. Med. 179:1029-1034

138. Tores BA. Griggs ND, Johnson HM. 1993. Bacterial and retroviral supermil-gent share a common binding region on class II MHC antigens. Nature 364:152-

McMuben CW, Bogazzki LX, Pullen AM. 1997. Mouse mammary runor virus su-perantigens require N-licked glycosylu-

perantigens require N-licked glycosylation for effective presentation to T colls. 
Notings 238.161-170
Labrecque N, Thibodess J, Mourad W, Schniy R-P. 1994. T cell receptor-major listocombathibity complex class II interaction is required for the T cell response to bacterial supermitigens. J. Exp. Med. 180:1921-1929
Deckhut AM, Chieo Y, Blackman MA, Woodland DL. 1994. Evidence for a fuer-tional internation however the II chain of

tional interaction between the fi chain of major histocompatibility complex clust II

major histocomparatinty empice cast it and the T cell receptor of chain during recognition of a bacterial superantigea, A. Exp. Med. 180:1931–1935
Blackman MA. Woodland DL. 1996. Role of the T cell receptor a-chain in superantigen recognition. Immunol. Res. 15:98–113

Pathm EA. 1994. Anatomy of the untihody molecule. Mol. Immunol. 31:169-

Fremont DH, Matsumura M, Sava EA, Peterson PA, Wilson IA. 1992. Crystal structures of two viral peptides in complex with MHC class I H-2Kb. Science 257:919-927

145. Madden D. 1995. The three-dimensional

145. Madden D. 1995. The three-dimensional structure of peptide-MHC complexes. Annu. Ros. Immunol. 13:587-622
146. Bhm TN, Bendley GA, Boulot G, Greche MI, Tello D, Doll'Acqua W, Bouchon H, Schwarz FP, Mariuzza RA, Poljak RI. 1994. Bound water molecules and conformational stabilization help mediate an andgen-antihody association. Proc. New. Acad. Sci. USA 91:1089-1093
147. Fields BA, Goldbaum FA, Ysern X, Poljak RJ, Maniuzza RA. 1995. Molecular basis of antigen minnery by an autidiotope. Nature 374:739-742
148. Dall'Acqua W, Goldman ER, Lin W, Teng C, Tsuchiya D, Li H, Ysern K, Braden BC. Li Y, Smith-Gill SJ, Mariuzza RA. 1998. A mutational analysis of binding inter-

A mutational analysis of binding inter-actions in an antigen-antibody proteinprotein complex. Biochamismy 37:7981-7991

Hodisov AS, Choi Y, Spanopoulou E, Pos-nett DN. 1998. My-orplastia superstriaca is a CDR3-dependent ligend for the T cell artigen receptor. J. Bap. Med. 187:319-327

Ciwrli C, Posnett DN, Schaly R-P, Denis F, 1998. Highly binsed CDR3 utage in restricted sets of \$6 chain variable regions during viral supermitiges 9 response. J. Exp. Med. 187:253-258
 Hong S-C, Waterbury G, Janeway CA Jr. 1996. Different supermitigens interact with distinct sites in the V\$6 domain of 0 single T cell receptor. J. Exp. Med. 183:143/-1446
 Davies DR, Padian EE. 1992. Taisting into chape. Carr. Biol. 2:254-256
 Tulip WR, Varghese JN, Laver WG, Wobster RG, Coleman PM. 1992. Refined crystal structure of the influenza virus N9 neuraminidate-NC41 Fab complex. J. Mol. Biol. 227:122-148
 Wilson JA, Statifield RL. 1993. Analgenantihody interactions. Carr. Opin. Struct.

antibody interactions. Curr. Opin. Struct.

Biol. 3:113-118

Lawrence MC, Coleman PM, 1993. Shape complementarity at protein protein interfaces. J. Mol. Biol. 234:946-950

156. Yearn X, Li H, Meriuzza RA. 1998. linperfect interfaces, Nature Struct. Biol.

- 5:412-414 157. Reich Z. Bonifice II, Lyons DS. Bono-chov N. Wachtel El, Davis MM. 1997. Ligand-specific oligometrication of T-cell receptor molecules. Nature 387:617-620
- 158. Imaaishi Kli, Igarashi H, Uchiyana T. 1990, Activation of murine T cells by streptococcal pyrogenic exotonin A. J. Immunol. 145:3170-3176

Immunol, 14513170-3170

159. Woodind DL, Blackman MA. 1993. How do T cell receptors. MBC molecules and superantigent get together? Immunol. Today 14:208-212

160. Barraro H, Donson D, Cervera C, Rexer C, Macphail S, 1995. T cell receptor Vada a extraorectal by a subsymmetric of VAS

is expressed by a subpopulation of VA6 T cells that respond to the basterial superantigen stophylosoccal enterotoxin B. J. Innunnal, 154:4247-4260

161. Donson D. Borrero H. Rutman M. Per-golizzi R. Mothado N. Maephall S. 1997. Gene transfer directly demonstrates a rote

- Gene transfer directly demonstrates rote for TCR Va elements in superantizen recognition. I. Immunol. 158:5229-5236
  Daly K., Nguyen P., Hankley D., Zhang WJ, Woodhand DL, Blackman MA. 1995.
  Contribution of the TCR a-chein to the differential recognition of besterial and extravial tunerantizens. I Immunol. and retrovirol superiolizeus. J. Immunol. 155:27-34
- Abrahmstn L. 1995. Superantigen engineering. Curr. Opin. Struct. Biol. 5:464-
- Dollaren M. Hedhud G. Akerbian E. Lando PA, Kalland T. 1991. Monocland

autibody-targeted superantigens: A dif-

ambody-targeted superantigens: A dif-ferent class of intil-tumor agents. Proc. Natl. Acad. Sci. USA 88:9287-9291

165. Doblisted M. Abrahmstin L. Bjöck P. Lando PA, Hedhund G, Forsberg G. Brodin T, Gascolome NRJ, Forberg C. Lind P, Katland T. 1994. Monoclound naibody-superantigen fixing proleins: Tumor-specific agents for T-cell-based temor therapy. Proc. Natl. Acad. Sci. USA 91:5945-8949

91:5945-3949

Ihle I, Holzer U, Krull F, Dohlsten M, Kelland T, Niethammer D, Damsecker GE, 1995. Antibody-targeted superantisens induce lysis of major histocompatibility complex class II-negative T-cell teukenia lines. Conver Res. 35:623-628

Holzer U, Bellage W, Krull F, Ihle J. Handgreinger R, Reisfeld RA, Dohlsten M, Kalland T, Niedhammer D, Danocker CE 1995. Superantisen-scaphylococcul-

in, enuano 1, recularante D, transcele GE. 1995. Superintigen-staphylococcul-enterotoxin-A-dependent and antihody-tergeted hyds of GD2-positive netwo-blasioma cells. Conver Immunol. Immu-nother. 41:129-136

notine: 41:129-436
Hanston J, Ohlsson L, Pereson R, Andersson G, Ilbiack N-G, Litton MJ, Kelland T, Dohlsten M, 1997. Genetically engineered superantigens as tolerable antitumor agents. Proc. Natl. Acad. Sci. USA

94:2489-2494

- 941.489-2494
  Giantonio Bl. Alpaugh RK, Schultz I, McAlect C, Newton DW, Shranton B, Guedez Y, Koth M, Vitek L, Perston R, Gunantson PG, Kalhand T, Dohlsten M, Persson B, Welner LM, 1997. Superantigen-based immunotherapy: a phase I stral of PNI-214565, a npy: a passe 1 trial of PNU-214565, a monetonal antibody-staphylocoscal enteroloxin A recombinant fusion prouch, in advanced panerciaic and colorectal cancer. J. Clin. Oncol. 15:1994-2007 Kepnier JW, Hetman A, Clements J, Marmek P. 1992. Mutations defining functional regions of the superantigen staphylococcal enterologia. R. J. Sep. 12-23
- lococcol enterotoxiu B. J. Asp. Med. 175:387-396 Brean S, Dyns B, Uirich RG, 1996.

Superscriptor vaccines: a comparative study of genetically attenuated receptor-binding mutants of studyly/lococcal emero-toxin A. J. Infact. Div. 174:338-345
Tsang I. Kamisas II., Trout RN, Huat RE, Chen IV, Johnson AJ, Piu I., Ruble DI. 1995. Humared immunity to acrosolized

sisphylococosi enterotoxin B (SEB), a fuprinningen, in monkeys exocimited with SEB toxeld-containing microspheres. In-foct. Immun. 68:2880-2885 Lowell GH, Colleton C, Frost D, Kaminski RW, Hughes M, Hatch J,

Hosper C, Estep J, Pitt L. Topper M, Hunt RE, Buker W, Bare WB, 1996. Immunogenicity and officacy against lethal acrosol staphylococcul enteroloxia B chollenge in mankeys by intramuscular and respiratory delivery of protecommunications. toxold vaccines. Infect. Immun. 64:4686-

McCormack JE, Callahan JE, Kappier J, Marrick PC, 1993. Profound deletion of mature T cells in vivo by chronic exposure

matture Teells in vivo by chronic exposure to exogenous superatilizen. J. Immunol. 150:3755-5792.

175. Michke T. Wahl C. Heeg K. Wagner H. 1993. Acquired resistance to superantigen-induced T cell shock. Vs telective T cell unresponsiveness unsides directly from a transient state of hyperceactivity. J. Immunol. 150:3776-3784.

176. Kim C. Simlnovitch KA, Ochi A. 1991. Reduction of lupus nephride in MRL/Ipr mice by a bacterial superantigen transment. J. Exp. Med. 174:1431-1437.

177. Rott O. Wekerle H. Fleicher B. 1992. Protection from experimental allergic encephalomycilis by application of a bacterial superantigen. Int. Immunol. 4:347-353.

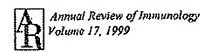
Soot IM, Schiffenbruer I, Johnson HM, 1993. Treatment of PLU mice with supermitigen staphytococcal enterotoxia B, prevents development of experimental allergic exceptalamyelitis. J. Neuroin-munal, 43:39-43

munal. 43:39-43
Racke MK, Quiglay L, Cannella B, Raine C, McFerlin DE, Scott DE. 1994. Supernuigen modulation of experimental allergic encephalomyclitis: activation of 
untagy determines outcome. J. Immunol. 152:2051-2059
Leung DYM, Schilevert PM. 1997. Superantigens in human disease. In Supernuigens: Molecular Biology. Immunology and Relavance to Human Disease, ed. 
DYM Leung, BT Huber, PM Schilevert, 
Marcel Dekker Int. NY, pp. 581-601
Hurley JM, Shintonkevitz R, Hanagan A, 
Eaney K, Boen E, Malmstron S, Kutzin 
BL, Matsumura M. 1995. Identification of 
class II major histocompatibility complex

Bl., Matsumura M. 1995. Identification of class II major histocompatibility complex and T cell receptor binding sites in the superantigen toxic shock syndrome toxin—1. J. Esp. Med. 181:2229-2235

Muray DL. Presed GS. Earhart CA. Leonard BAB, Kreiswirds BN, Novick RP, Ohlendorf DH, Schlievert PM. 1994, immunolishogie and hischemical properties of minutes of straight syndrome.

immunoushogic and hischemical proper-tice of mutuals of toxic shock syndrome toxin-1. J. Immunol. 152:37-95 Derostewicz RL, Word J, Chan M, Fin-berg RW, Kasper DL. 1994. Mutations affecting the activity of toxic shock syn-drome toxin-1. Biochemistry 33:17844-12851



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